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54) Title: METHOD TO IDENTIFY COMPOUNDS FOR	DISRI	PTING PROTEIN/PROTEIN INTERACTION	ıs
57) Abstract	•	~ .	
The present invention relates generally to materials and methods for identification of inhibitors of interactions etween known binding partner proteins.	_	Y Pres	
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# METHODS TO IDENTIFY COMPOUNDS FOR DISRUPTING PROTEIN/PROTEIN INTERACTIONS

#### Background of the Invention

The present invention relates to a novel method to identify inhibitors of protein/protein interactions.

#### Background

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Modulation of protein/protein interactions is an attractive target for drug discovery and development. Potential methods by which drugs can regulate protein/protein interactions are numerous, including, for example, regulation of expression of one or more of the binding proteins, modulation of post-translational modification, and direct interference with the capacity of one protein to bind to one or more binding partners. More importantly, recent observations make it increasingly clear that supramolecular protein complexes, involving two or more binding proteins, play an important and essential roles in signal transduction, gene expression, cell proliferation and duplication, and cell cycle progression. For example, in the repair of UV damaged DNA, a so-called "repairsome" that contains over ten individual proteins is assembled into a complex which can then carry out the necessary repair. Likewise, gene transcription occurs through the concerted action of greater than twenty proteins. Signal transduction proteins, such as receptor protein kinases, are part of large complexes with many proteins. Contacts through Src homology type 2 (SH2) domains on the receptor kinases, for example, are noteworthy protein interaction which are part of one or more enzymatic cascade important for many metabolic processes. Disrupting the binding capacity of one or more proteins which form any of these larger complex is therefore an important and untapped method to control action of the overall complex.

Protein/protein interactions have been discovered and characterized by a variety of methods: (i) standard biochemical affinity

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methods such as chromatography or co-immunoprecipitations; (ii) gel overlay methods; (iii) co-purification by traditional biochemistry; and (iv) two-hybrid analysis [Fields and Song, Nature 340:245-246 (1989); Fields, Methods: A Companion to Methods in Enzymology 5:116-124 (1993); U.S. Patent 5, 283, 173 issued February 1, 1994 to Fields, et al.]. The most recent of these approaches, the two hybrid method, has enjoyed broad application because of its relative ease of use for gene identification from cDNA fusion libraries. [See Chien et al., Proc. Natl. Acad. Sci. (USA) 88:9578-9582 (1991); Dalton and Treisman, Cell 72:223-232 (1993); and Durfee, et al., Genes and Devel. 7:555-569 (1993)].

The two hybrid system is based on targeting and identifying a protein/protein interaction through the use of a reporter system. The described two hybrid systems either use the yeast Gal4 DNA binding domain or the E. coll lexA DNA binding domain and couple this region to a transcriptional activator such as Gal4 or VP16 that drives a reporter like  $\beta$  galactosidase or HIS3.

In principle the two hybrid assay could be used for drug screening. [See WO 96/03501 and WO 96/03499.] In such a scenario, loss of  $\beta$  galactosidase or HIS3 activity would be identified after the yeast strain is treated with a compound. In practice, however, use of the two hybrid system is technically undesirable for several reasons. In instances where the  $\beta$  galactosidase or HIS3 protein are employed as the reporter protein, a loss of activity is particularly difficult to detect because the expressed reporter protein is too long lived to be used in a high throughput mode. If a candidate binding inhibitor compound is metabolized faster than the previously expressed reporter protein is turned over, it is difficult to detect inhibitory action of the candidate drug while a reporter protein is still active. In high throughput screening, the loss of a positive signal, for example,  $\beta$  galactosidase or HIS3 is impossible to detect. Present robotocized screening and detection methods are simply not sufficiently sensitive or robust to detect loss of a signal.

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Thus there is a need in the art to develop a rapid screening method that gives a positive signal, as opposed to a negative signal, when a protein/protein interaction is disrupted. Such a system must be capable of using protein interactions that are initially detected by any of the above mentioned approaches and must be sufficiently robust to detect a gain of function when a protein interaction is lost. In essence, the screening method must give a signal when an interaction is lost, not lose a signal when an interaction is lost. Such a system must be sensitive to subtle interactions, in particular ones that are caused by post-translational modification like protein phosphorylation. Finally for large scale screening, such as high throughput screening, the system must be manipulable such that a large signal-to-noise ration can be easily detected.

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#### Brief Summary of the Invention

In one aspect, the present invention provides materials that are useful for the identification of compounds which inhibit interaction between known binding partner proteins. See Figure 1. The invention provides host cells transformed or transfected with DNA comprising: (i) a repressor gene encoding DNA binding protein that acts as a repressor protein, said repressor gene under transcriptional control of a promoter; (ii) a selectable marker gene encoding a selectable\marker protein; said selectable marker gene under transcriptional control of an operator; said operator regulated by interaction with said repressor protein; (iii) a first recombinant fusion protein gene encoding a first binding protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activating protein or a transactivating domain of a transcriptional activating protein; and (iv) a second recombinant fusion protein gene encoding a second binding protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activating protein or a transactivating domain of a transcriptional activating protein, whichever domain is not encoded by the first

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fusion protein gene, said second binding protein or binding fragment thereof capable of interacting with said first binding protein or binding fragment thereof such that interaction of said second binding protein or binding fragment thereof and said first binding protein or binding fragment thereof brings into proximity a DNA binding domain and a transactivating domain forming a functional transcriptional activating protein; said functional transcriptional activating protein or said promoter to increase expression of said repressor sene.

The invention comprehends host cells wherein the various genes and regulatory sequences are encoded on a single DNA molecule as well as host cells wherein one or more of the repressor gene, the selectable marker gene, the first recombinant fusion protein gene, and the second recombinant fusion protein gene are encoded on distinct DNA expression constructs. In a preferred embodiment, the host cells are transformed or transfected with DNA encoding the repressor gene, the selectable marker gene, the first recombinant fusion protein gene, and the second recombinant fusion protein gene, each encoded on a distinct expression construct. Regardless of the number of DNA expression constructs introduced, each transformed or transfected DNA expression construct further comprises a selectable marker gene sequence, the expression of which is used to confirm that transfection or transformation was, in fact, accomplished. Selectable marker genes encoded on individually transformed or transfected DNA expression constructs are distinguishable from the selectable marker under transcriptional regulation of the tet operator in that expression of the selectable marker gene regulated by the tet operator is central to the preferred embodiment; i.e., regulated expression of the selectable marker gene by the tet operator provides a measurable phenotypic change in the host cell that is used to identify a binding protein inhibitor. Selectable marker genes encoded on individually transformed or transfected DNA expression constructs are provided as determinants of successful transfection or transformation of the individual

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DNA expression constructs. Preferred host cells of the invention include transformed S. cerevisiae strains designated Y1596 and Y1584 which were deposited August 13, 1996 with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland 20852, and assigned Accession Numbers ATCC 74384 and ATCC 74385, respectively.

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The host cells of the invention include any cell type capable of expressing the heterologous proteins required as described above and which are capable of being transformed or transfected with functional promoter and operator sequences which regulate expression of the heterologous proteins also as described. In a preferred embodiment, the host cells are of either manmal, insect or yeast origin. Presently, the most preferred host cell is a yeast cell. The preferred yeast cells of the invention can be selected from various strains, including the S. cerevisiae yeast transformants described in Table 1. Alternative yeast specimens include S.pombe, K.lactis, P.pastoris, S.carlsbergensis and C.albicans. Preferred mammalian host cells of the invention include Chinese hamster ovary (CHO), COS, HeLa, 3T3, CV1, LTK, 293T3, Rat1, PC12 or any other transfectable cell line of human or rodent origin. Preferred insect cells lines include SF9 cells.

In a preferred embodiment, the selectable marker gene is regulated by an operator and encodes an enzyme in a pathway for synthesis of a nutritional requirement for said host cell such that expression of said selectable marker protein is required for growth of said host cell on media lacking said nutritional requirement. Thus, as in a preferred embodiment where a repressor protein interacts with the operator, transcription of the selectable marker gene is down-regulated and the host cells are identified by an inability to grow on media lacking the nutritional requirement and an ability to grow on media containing the nutritional requirement. In a most preferred embodiment, the selectable marker gene encodes the HIS3 protein, and host cells transformed or transfected with a HIS3-encoding DNA expression construct are selected following growth on media in the presence

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and absence of histidine. The invention, however, comprehends any of a number of alternative selectable marker genes regulated by an operator. Gene alternatives include, for example URA3, LEU2, LYS2 or those encoding any of the multitude of enzymes required in various pathways for production of a nutritional requirement which can be definitively excluded from the media of growth. In addition, conventional reporter genes such as chloramphenicol acetyltransferase (CAT), firefly luciferase, β-galactosidase (β-gal), secreted alkaline phosphatase (SEAP), green fluorescent protein (GFP), human growth hormone (hGH), β-glucuronidase, neomycin, hygromycin, thymidine kinase (TK) and the like may be utilized in the invention.

In the preferred embodiment, the host cells include a repressor protein gene encoding the tetracycline resistance protein which acts on the tet operator to decrease expression of the selectable marker gene. The invention, however, also encompasses alternatives to the tet repressor and operator, for example, E. coli trp repressor and operator, his repressor and operator, and lac operon repressor and operator.

The DNA binding domain and transactivating domain components of the fusion protein may be derived from the same transcription factor or from different transcription factors as long as bringing the two domains into proximity permits formation of a functional transcriptional activity protein that increases expression of the repressor protein with high efficiency. A high efficiency transcriptional activating protein is defined as having both a DNA binding domain exhibiting high affinity binding for the recognized promoter sequence and a transactivating domain having high affinity binding for transcriptional machinery proteins required to express repressor gene mRNA. The DNA binding domain component of a fusion protein of the invention can be derived from any of a number of different proteins including, for example, LexA or Gal4. Similarly, the transactivating component of the invention's fusion proteins can be derived from a number of different transcriptional activating proteins, including for example, Gal4 or

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VP16. In one embodiment of the invention, polynucleotides encoding binding partner proteins CREB and CBD are inserted in plasmids pVP16-CREB and pLexA-CBD, respectively, which were deposited with the ATCC and assigned Accession Numbers ATCC 98138 and ATCC 98139, respectively.

The promoter sequence of the invention which regulates transcription of the repressor protein can be any sequence capable of driving transcription in the chosen host cell. The promoter may be a DNA sequence specifically recognized by the chosen DNA binding domain of the invention, or any other DNA sequence with which the DNA binding domain of the fusion protein is capable of high affinity interaction. In a preferred embodiment of the invention, the promoter sequence of the invention is either a HIS3 or alcohol dehydrogenase (ADH) promoter. In a presently most preferred embodiment, the ADH promotor is employed in the invention. The invention, however, encompasses numerous alternative promoters, including, for example, those derived from genes encoding HIS3, ADH, URA3, LEU2 and the like.

In another aspect, the invention provides methods to identify molecules that inhibit interaction between known binding partner proteins. In one embodiment, the invention provides a method to identify an inhibitor of binding between a first binding protein or binding fragment thereof and a second binding protein or binding fragment thereof comprising the steps of (a) growing host cells transformed or transfected as described above in the absence of a test compound and under conditions which permit expression of said first binding protein or binding fragment thereof and said second binding protein or binding fragment that said first binding protein or fragment thereof and second binding protein or binding fragment thereof interact bringing into proximity said DNA binding domain and said transactivating domain forming a functional transcriptional activating protein; the transcriptional activating protein acting on said promoter to increase

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expression of said repressor protein; said repressor protein interacting with said operator such that said selectable marker protein is not expressed; (b) confirming lack of expression of said selectable marker protein in said host cell; (c) growing said host cells in the presence of a test compound; and (d) comparing expression of said selectable marker protein in the presence and absence of said test compound wherein increased expression of said selectable marker protein is indicative that the test compound is an inhibitor of binding between said first binding protein or binding fragment thereof and said second binding protein or binding fragment thereof.

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In a most preferred embodiment, the invention provides a method to identify an inhibitor of binding between a first binding protein or binding fragment thereof and a second binding protein or binding fragment thereof comprising the steps of: (a) transforming or transfecting a host cell with a first DNA expression construct comprising a first selectable marker gene encoding a first selectable marker protein and a repressor gene encoding a repressor protein, said repressor gene under transcriptional control of a promoter; (b) transforming or transfecting said host cell with a second DNA expression construct comprising a second selectable marker gene encoding a second selectable marker protein and a third selectable marker gene encoding a third selectable marker protein, said third selectable marker gene under transcriptional control of an operator, said operator specifically acted upon by said repressor protein such that interaction of said repressor protein with said operator decreases expression of said third selectable marker protein; (c) transforming or transfecting said host cell with a third DNA expression construct comprising a fourth selectable marker gene encoding a fourth selectable marker protein and a first fusion protein gene encoding a first binding protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activation protein or a transactivating domain of said transcriptional activation protein; (d) transforming or transfecting said host cell with a fourth DNA expression construct comprising

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a fifth selectable marker gene encoding a fifth selectable marker protein and a second fusion protein gene encoding a second binding protein or binding fragment thereof in frame with either the DNA binding domain of said transcriptional activation protein or the transactivating domain of said transcriptional activation protein, whichever is not included in first fusion protein gene; (e) growing said host cell under conditions which permit expression of said first binding protein or fragment thereof and said second binding protein or fragment thereof such that said first binding protein or fragment thereof and second binding protein or binding fragment thereof interact bringing into proximity said DNA binding domain and said transactivating domain reconstituting said transcriptional activating protein; said transcriptional activating protein acting on said promoter to increase expression of said repressor protein; said repressor protein interacting with said operator such that said third selectable marker protein is not expressed; (f) detecting absence of expression of said selectable gene; (g) growing said host cell in the presence of a test compound of binding between said first protein or fragment thereof and said second binding protein or fragment thereof; and (h) comparing expression of said selectable marker protein in the presence and absence of said test compound wherein decreased expression of said selectable marker protein is indicative of an ability of the test compound to inhibit binding between said first binding protein or binding fragment thereof and said second binding protein or binding fragment thereof such that said transcriptional activating protein is not reconstituted, expression of said repressor protein is not increased, and said operator increases expression of said selectable marker protein.

The methods of the invention encompass any and all of the variations in host cells as described above. In particular, the invention encompasses a method wherein: the host cell is a yeast cell; the selectable marker gene encodes HIS3; transcription of the selectable marker gene is regulated by the *tet* operator; the repressor protein gene encodes the

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tetracycline resistance protein; transcription of the tetracycline resistance protein is regulated by the HIS3 promoter; the DNA binding domain is derived from LexA; and the transactivating domain is derived from VP16. In another embodiment, the invention encompasses a method wherein: the lost cell is a yeast cell; the selectable marker gene encodes HIS3; transcription of the selectable marker gene is regulated by the *tet* operator; the repressor protein gene encodes the tetracycline resistance protein; transcription of the tetracycline resistance protein is regulated by the alcohol dehydrogenase promoter; the DNA binding domain is derived from LexA; and the transactivating domain is derived from VP16.

In alternative embodiments of the invention wherein the host cell is a mammalian cell, variations include the use of mammalian DNA expression constructs to encode the first and second recombinant fusion genes, the repressor gene, and the selectable marker gene, and use of selectable marker genes encoding antibiotic or drug resistance markers (i.e., neomycin, hygromycin, thymidine kinase).

There are at least three different types of libraries used for the identification of small molecule modulators. These include: (1) chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of random peptides, oligonucleotides or organic molecules.

Chemical libraries consist of structural analogs of known compounds or compounds that are identified as "hits" via natural product screening. Natural product libraries are collections of microorganisms, animals plants or marine organisms which are used to create mixtures for screening by: (1) fermentation and extraction of broths from soil, plant or marine microorganisms or (2) extraction of plants or marine organisms. Combinatorial libraries are composed of large numbers of peptides, oligonucleotides or organic compounds as a mixture. They are relatively easy to prepare by traditional automated synthesis methods, PCR, cloning or proprietary synthetic methods. Of particular interest are peptide and

oligonucleotide combinatorial libraries. Still other libraries of interest include peptide, protein, peptidomimetic, multiparallel synthetic collection, recombinatorial, polypeptide libraries.

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The utility of the various aspects of the invention is manifest. Host cells of the invention are useful to demonstrate *in vivo* binding capacity of both known and suspected binding partner proteins in a recombinant system. Such an expression system permits systematic analysis of the structure and function of a particular binding protein, thus permitting identification and/or synthesis of potential modulators of the physiological activity of the binding proteins. The methods of the invention are particularly useful to identify and improve molecules which are capable of inhibiting specific and general protein/protein interactions. Inhibitors identified by the methods of the invention can then be examined for utility *in vivo* as therapeutic and/or prophylactic medicaments for conditions associated with various protein/protein interactions.

#### Description of the Drawing

Figure 1 describes the mechanics of the split hybrid assays.

### Detailed Description of the Invention

The present invention relates generally to methods designated split hybrid assays to identify inhibitors of protein/protein interactions and is illustrated by the following examples describing various methods for making and using the invention. In particular, Example 1 relates to construction of various plasmids and expression constructs utilized in the invention. Example 2 described generation of various yeast transformants used to identify inhibitor compounds. Examples 3, 4, 5 and 6 address use of the split hybrid assay to examine CREB/CBD binding, Tax/SRF binding, CKI/CREB binding and AKAP 79 binding to various partner protein, respectively. Example 7 describe general application of the split hybrid assay. Example 8 relates to

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use of the split hybrid assay for weakly interacting binding partners. Example 9 describes general assay methods. Example 10 addresses use of the split hybrids assay to identify agents that prevent receptor desensitization and drug tachyphylaxis.

#### Example 1 Plasmid Construction

In the examples that follow, various plasmid constructs were utilized as described. To simplify discussion of the exemplified assays, this example describes construction of the various plasmids used in the following examples. For clarity, the plasmids are grouped according common features relating to their applications in the assays later discussed.

#### I. Plasmids Encoding Reporter Gene HIS3

#### A. pRS303/1xtetop-MluI

One copy of the tet operator sequence was engineered into position -53 in the HIS3 promoter of pRS313 [Sikorski, R.S. et al., Genetics 122:19-27 (1989)] by using the polymerase chain reaction (PCR). Two primary PCR reactions using pRS313 as a template were performed which utilized a 5'-terminal oligonucleotide designated Eco47III-5' and a 3'-inner oligonucleotide designated Tetop internal 3' to yield a primary 5'-PCR product and a 5'-inner oligonucleotide designated Tetop internal 5 and a 3'-terminal oligonucleotide designated Nhe I 3' to yield a primary 3'-PCR product.

Eco47 III-5' SEQ ID NO: 1 5'-TTGGTGAGCGCTAGGAGTCACTGCCAG

Tetop int. 3' SEQ ID NO: 2

5'-TATACTCTATCAATGATAGAGTAATTCATTATGTGATAATGCC

Tetop int. 5' SEQ ID NO: 3
5'-ATTACTCTATCATTGATAGAGTATATAAAGTAATGTGATTTC)

Nhe I 3'

SEO ID NO: 4

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#### 5'-AATTCTGCTAGCCTCTGCAAAGC

5' and 3' inner oligonucleotides contain complementary sequence such that 3' sequence of the primary 5' PCR product overlaps with 5' sequence of the primary 3' PCR product. The 5' terminal oligonucleotide contains the restriction site Eco47III while the 3' terminal oligonucleotide contains the 5 restriction site NheI in order to facilitate subsequent subcloning. The primary PCR reactions were performed with Pfu DNA polymerase (Stratagene, La Jolla, CA) using reaction conditions described by the manufacturer. PCR products were isolated by Bio101 (Vista, CA) Gene Clean III gel extraction. 10 The primary 5' and 3' PCR products were then combined in a second PCR reaction and amplified using the 5'- and 3'- terminal oligonucleotides. Eco47III-5' and Nhe I 3'. The second PCR reaction was performed with Vent DNA polymerase (New England Biolabs, Beverly, MA) using reaction conditions described by the manufacturer, except that the reactions were 15 supplemented with 4 mM Mg2+. The final PCR product contained one tet operator sequence inserted into position -53 of the HIS3 promoter and nucleotides 52-48 deleted in the construction. The final PCR product was isolated, digested with Eco47III and NheI and cloned into pRS313 previously digested with Eco47III and NheI. The resulting plasmid was designated pRS313/1xtetop. DNA sequencing confirmed the presence of one copy of the 20 tet operator sequence in pRS313/1xtetop and confirmed integrity of the Eco47III and NheI junctions

A Mlul restriction enzyme site was engineered into position -22 in the HIS3 promoter of pRS313/1xtetop by utilizing PCR using Vent DNA polymerase using pRS313/1xtetop as template. One PCR construct was amplified using the 5' terminal oligonucleotide Eco47 III-5' (SEQ ID NO: 1) containing an Eco47III restriction site and a 3'-oligonucleotide designated Mlu I 3' containing a Mlul restriction site.

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## Mlu I 3' SEQ ID NO: 5 5'-CGCACGCGTCGAAGAAATCACATTACTTTATATA

A second PCR product was amplified using the 3'-terminal oligonucleotide Nhe I 3' (SEQ ID NO: 4) containing a NheI restriction site and a 5'oligonucleotide designated Mlu I 5' containing a MluI restriction site.

## Mlu I 5' SEQ ID NO: 6 5'-CGCACGCGTATACTAAAAAATGAGCAGGCAAG

The first PCR product was isolated and digested with Eco47III and MluI, while the second PCR product was isolated and digested with MluI and NheI. These digested products were isolated and ligated in a triple ligation with pRS313 previously digested with Eco47III and NheI. The resulting plasmid we designated pRS313/1xtetop-MluI. DNA sequencing confirmed the presence of the MluI site in pRS313/1xtetop-MluI and confirmed that integrity of the Eco47III and NheI junctions were maintained.

A pRS303/1xtetop-Mlul plasmid was constructed by first removing the Eco47III/Nhel fragment containing the altered HIS3 promoter from the pRS313/1xtetop-Mlul vector and ligating the isolated fragment into pRS303 previously digested with Eco47III and Nhel. DNA sequencing confirmed proper insertion of the Eco47IIII/Nhel fragment.

#### B. pRS303/2xtetop-LYS2

One copy each of the tet operator sequence was engineered into positions -53 and -22 in the HIS3 promoter of pRS303 (Sikorski, et al., Genetics 122:19-27 (1989)). PCR was utilized to engineer one copy into position -53 which resulted in plasmid pRS303/1xtetop. To insert the second copy, a Mlul site was introduced at position -22 in the HIS3 promoter using PCR. The new plasmid was designated pRS303/1xtetop-MluI.

The tet operator was created by annealing two complementary

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oligonucleotides tetop-1 and tetop-2.

tetop-1 SEQ ID NO: 7
5'-CGCGTACTCTATCATTGATAGAGTA;

tetop-2 SEQ ID NO: 8
5'-ATGAGATAGTAACTATCTCATGCGC

When annealed, the tet operator sequence contains flanking Mtul sites. Both oligonucleotides were phosphorylated using 7d polynucleotide kinase (Gibco BRL, Grand Island, NY) at 37°C for one hour and annealed by first heating at 70°C for 10 minutes and then cooling to room temperature. The annealed oligonucleotides were isolated and ligated into pRS303/1xtetop-Mtul previously digested with Mtul. The resulting plasmid was designated pRS303/2xtetop. DNA sequencing confirmed insertion of one copy of the tet operator sequence in the Mtul site.

The LYS2 gene was digested from pLYS2 [Hollenberg, S.M. et al., Mol. Cell.Biol. 15:3813-3822 (1995)] with EcoRI and HindIII and the isolated fragment blunt ended using the large fragment of DNA polymerase I (Gibco BRL, Grand Island, NY). Phosphorylated SxII linkers (New England Biolabs, Beverly, MA) were ligated to the fragment, the fragment digested with SxII, and the resulting fragment ligated into pRS313 previously digested with SxII. The resulting plasmid was designated pRS313/LYS2.

The LYS2 fragment was removed from pRS313/LYS2 with Ssrl digestion and inserted into pRS303/2xtetop previously digested with Ssrl. The resulting plasmid was designated pRS303/2xtetop-LYS2.

Similarly, the LYS2 SxiI fragment was inserted into pRS303/1xtetop-MluI previously digested with SxiI yield pRS303/1xtetop-MluI-LYS2.

#### C. pRS303/3xtetop-LYS2

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Two copies of the tet operator sequence were created by selfannealing a palindromic oligonucleotide Tetop 2x with itself.

## Tetop 2x SEQ ID NO: 9 5'-CGCGTACTCTATCATTGATAGAGTCTAGACTCTATCAATGATAGAGTA

5 The annealed oligonucleotide contained flanking MluI sites. The oligonucleotide was phosphorylated, annealed, and isolated as above. The isolated annealed and MluI-digested oligonucleotide was ligated into pRS303/1xtetop-MluI-LYS2 previously digested with MluI to yield pRS303/3xtetop-LYS2. The presence of two copies of the tet operator 10 sequence in the MluI site was confirmed by DNA sequencing.

#### D. pRS303/4xtetop-LYS2 and pRS303/8xtetop-LYS2

Three or seven copies of the *ter* operator were created using PCR with *Vent* DNA polymerase as described above. Plasmid pUHC-13-3 [Grossen and Bujarg, *Proc. Natl. Acad. Sci. (USA)* 82:5547-5551 (1992)] was nsed as template DNA using 5'- and 3'- oligonucleotides, Mlu I/Sph I 5' and Mlu I Sph I 3', containing an exterior *MluI* restriction enzyme site nested internally by a *SphI* restriction enzyme site.

Mlu I/Sph 1 5' SEQ ID NO: 10
5'-GCGACGCGTGCATGCCGTCTTCAAGAATTCCTCGAG

20 Mlu I Sph I 3' SEQ ID NO: 11 5'-GCGACGCGTGCATGCCCACCGTACACGCCTACTCGA

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The PCR products were separated on an agarose gel and the ladder of different sized DNA fragments was isolated, digested with MluI, and ligated into the MluI restriction site of pRS303/1xtetop-MluI-LYS2. DNA sequencing revealed that either three or seven copies of tet operators were inserted into the Mlu site of pRS303/1xtetop-MluI-LYS2 to provide either pRS303/4xtetop-LYS2 or pRS303/8xtetop-LYS2.

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### E. pRS303/6xtetop-LYS2 and pRS303/10xtetop-LYS2

A SphI restriction enzyme site was introduced at position -85 in the HIS3 promoter of pRS303/3xtetop-LYS2 using PCR with Vent DNA polymerase as described. Plasmid pRS303/3xtetop-LYS2 was used as a template DNA. A first fragment was amplified using the 5'-terminal oligonucleotide Eco47 III-5' (SEQ ID NO: 1) described above containing an Eco47III restriction site and a 3'-oligonucleotide Sph I 3' containing a SphI restriction site.

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Sph I 3' SEQ ID NO: 12
5'-CATGGCATGCAAAAAAAAAAGGGTCATCCGCTAGG

A second PCR product was amplified using the 3'-terminal oligonucleotide Nhe I 3' (SEQ ID NO: 4) described above containing a NheI restriction site and a 5'-oligonucleotide containing a SphI restriction site.

Sph I 5' SEQ ID NO: 13 5'CATGGCATGCTTAGCGATTGGCATTATCACAT

The PCR products were isolated as described above. The first PCR product was digested with Eco47III and SphI, and the second PCR product was digested with SphI and NheI. Both digestion products were ligated in a triple ligation along with pRS303/3xtetop-LYS2 previously digested with both Eco47III and NheI. The resulting plasmid was designated pRS303/3xtetop-SphI-LYS2. The presence of the SphI site in pRS303/3xtetop-SphI-LYS2 was confirmed by DNA sequencing analysis.

Three copies of tet operators were isolated as a single fragment by digesting pRS303/4xtetop-LYS2 with SphI. The isolated fragment was ligated into the SphI site of pRS303/3xtetop-SphI-LYS2 to yield pRS303/6xtetop-LYS2. The presence of three additional copies of the tet operator in pRS303/6xtyetop-LYS2 at the SphI site was confirmed by DNA

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sequencing.

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Seven copies of *tet* operators were isolated as a single fragment by digesting pRS303/8xtetop-LYS2 with *SphI*. The isolated fragment was ligated into the *SphI* site of pRS303/3xtetop-*SphI*.LYS2 to yield pRS303/10xtetop-LYS2. The presence of seven additional copies of the tet operator in pRS303/10xtetop-LYS2 at the *SphI* site was confirmed by DNA sequencing.

#### F. pRS313/MluI and pRS303/MluI

A MluI restriction enzyme site was engineered into position -22 in the HIS3 promoter of pRS313 utilizing PCR and Vent DNA polymerase as noted above. Plasmid pRS313 was used as a template for these PCR One PCR construct was amplified using the 5' terminal oligonucleotide Eco47 III-5' (SEQ ID NO: 1) containing an Eco47III restriction site and a 3' oligonucleotide Mlu I 3' (SEQ ID NO: 5) containing a MluI restriction site. A second PCR product was amplified using the 3' terminal oligonucleotide Nhe I 3' (SEQ ID NO: 4) containing a NheI restriction site and the 5' oligonucleotide Mlu I 5' (SEQ ID NO: 6) containing a Mlul restriction site. The first PCR product was isolated and digested with Eco47III and MluI, while the second PCR product was isolated and digested with MluI and NheI. The digested products were partially purified and joined in a triple ligation with pRS313 which had been previously digested with Eco47III and Nhel. The resulting plasmid was designated pRS313/MluI. DNA sequencing confirmed the presence of the Mlul site in pRS313/Mlul and to confirm the integrity of the Eco47III and NheI junctions.

pRS303/MluI was constructed in exactly the same manner as pRS313/MluI except that pRS303 was used in place of pRS313.

#### G. pRS313/1xtetop

See above wherein pRS313/1xtetop is an intermediate in the

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construction of pRS303/1xtetop-MluI.

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#### H. pRS313/MluI-1xtetop and pRS303/MluI-1xtetop

One copy of the *tet* operator sequence was created by annealing two complementary oligonucleotides tetop-1 and tetop-2 (SEQ ID NO: 7 and SEQ ID NO: 8). The annealed *tet* operator sequence contains flanking *Mlul* sites. The oligonucleotides were phosphorylated using *T4* polynucleotide kinase (Gibco BRL, Grand Island, NY) at 37°C for one hour and annealed by first heating at 70°C for 10 minutes followed by cooling to room temperature. The annealed oligonucleotides were isolated and ligated separately into *Mlul*-digested pRS313/Mlul and pRS303/Mlul, the resulting plasmids being designated pRS313/Mlul-1xtetop and pRS303/Mlul-1xtetop. DNA sequencing confirmed the presence of one copy of the *tet* operator in the *Mlul* sites of both plasmids.

In order to produce plasmids bearing multiple copies of the tet operator, annealed oligonucleotides described above were ligated together overnight at 16°C. After isolation of the ligation products, they were inserted into the Mlul of pRS313/Mlul. DNA sequencing analysis confirmed that one clone, pRS313/Mlul-4xtetop, was produced which contained four copies of tet operator in the Mlul site. However, upon further examination of this clone it was discovered that it had been subjected to a recombination event and was therefore not useful for further cloning steps. Continued attempts to insert multiple copies of the tet operator into the Mlul site of pRS313/Mlul by ligating multimers of the tet operator have been unsuccessful.

#### pRS313/1xtetop-MluI

See above wherein construction of pRS313/1xtetop-MluI was an intermediate in the construction of pRS303/1xtetop-MluI.

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#### J. pRS313/2xtetop

One copy of the *tet* operator sequence was created using annealed complementary oligonucleotides tetop-1 and tetop-2 (SEQ ID NO: 7 and SEQ ID NO: 8). Annealed oligonucleotides were ligated into the *Mlul* site of pRS313/1xtetop-Mlul to yield pRS313/2xtetop. DNA sequencing confirmed the presence of two copies of the *tet* operator in the *Mlul* site.

#### K. pRS303/2xtetop

See above wherein pRS303/2xtetop was an intermediate in the construction of pRS303/2x/tetop-LYS2.

### L. pRS313/LYS2 and pRS313/LYS2

The LYS2 gene was digested from pLYS2 with EcoRI and HindIII digestion. The EcoRI/HindIII fragment was blunt ended using the large fragment of DNA polymerase I (Gibco BRL, Grand Island, NY) and ligated with phosphorylated SxII linkers (New England Biolabs, Beverty, MA).

15 The resulting fragment was digested with SxII and ligated into pRS313 previously digested with SxII. The resulting plasmid was designated pRS313/LYS2. Because the LYS2 fragment was shown to have inserted into pRS313 in both orientations, plasmids with the LYS2 gene in both orientations were transformed separately into the yeast strain SEY6210α\_(MATα\_leu2-20] 3.112 ura3-52 his3-Δ200 trp1-Δ901 lys2-801 suc2-Δ9 [Robinson et al., Mol. Cell. Biol. 8:4936-4948 (1988)]. Both clones allowed the yeast to grow in the absence of lysine indicating that orientation of the LYS2 gene in pRS313 did not affect the expression of an active gene.

The LYS2 fragment was removed from pRS313/LYS2 with Sx12
25 and ligated into the Sx12 site of:

pRS313/1xtetop-MluI giving plasmid pRS313/1xtetop-MluI-LYS2, pRS313/2xtetop giving plasmid pRS313/2xtetop-LYS2,

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pRS303/1xtetop-MluI giving plasmid pRS303/1xtetop-MluI-LYS2, and pRS303/2xtetop giving plasmid pRS303/2xtetop-LYS2.

#### II. Plasmids Encoding Reporter Gene TetR

#### A. pRS306/HIS3:TetR/Term

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The 5' promoter sequence of the yeast HIS3 gene, encompassing nucleotides -75 to +23, was ligated to the translational start of TetR. In addition, the DNA sequence encoding the simian virus 40 (SV40) large T antigen nuclear localization signal was ligated in frame with the nucleotide sequence encoding the last amino acid residue of TetR. The chimeric fragment was created by the same PCR strategy as described above.

The HIS3 promoter fragment, the primary 5'-PCR product, was amplified by PCR from plasmid p601 [Grueneberg, D.A., Science 257:1089-1095 (1992)] using a 5'-terminal oligonucleotide T7 Promoter primer and a 3'-inner oligonucleotide 3'-TetR inner primer.

T7 Promoter primer SEQ ID NO: 14 5'-TAATACGACTCACTATATAGGG

> 3'-TetR inner primer SEQ ID NO: 15 5'-TCTAGACTTTGCCTTCGTTTATC

The primary 3' PCR product containing the *TetR* coding sequence was amplified from pSLF104 [Forsburg, *Nucl. Acid. Res.* 21:2955-2956 (1993)] with a 5'-inner oligonucleotide 5'-TetR inner primer and a 3'-terminal oligonucleotide 3'-TetR terminal primer.

5'-TetR inner primer SEQ ID NO: 16 5'-CGAAGGCAAAGATGTCTAGATTAGATAAAAG

3'-TetR terminal primer SEQ ID NO: 17
5'-CGCGGATCCGCTTTCTCTTTTTTGGAGACCCACTTTCACATTTAAG

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An EcoRI site derived from the p601 fragment and a BamHI site in the 3'terminal oligonucleotide were used in subsequent subcloning. The PCR
products were gel-purified and amplified in a second PCR reaction with 5'and 3-' terminal oligonucleotides, T7 Promoter primer (SEQ ID NO: 14) and
3'-TetR terminal primer (SEQ ID NO: 17). The secondary PCR product was
isolated, digested with EcoRI and BamHI, and ligated into pR\$306/Term
previously digested with EcoRI and BamHI. The resulting plasmid was
designated pR\$306/HIS3-TetR/Term which comprises the complete TetR
coding sequence in frame with sequences encoding the nuclear localization
signal of SV40 large T antigen.

#### B. pRS316/HIS3:TetR/Term

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The construction protocol for this plasmid was the same as described above for subcloning a HIS3 DNA into pRS306/Term except that the vector for subcloning was pRS316/Term described above.

#### C. pRS306/1xLexAop/HIS3:TetR

Oligonucleotides LexAop (100a) and LexAop (100b) containing a single copy of LexA operator were phosphorylated with 74 polynucleotide kinase (Gibco BRL, Grand Island, NY) at 37°C for one hour.

LexAop (100a) SEQ ID NO: 18 5'-AATTGCTCGAGTACTGTATGTACATACAGTAG

LexAop (100b) SEQ ID NO: 19 5'-AATTCTACTGTATGTACATACAGTACTCGAGC

Following phosphorylation, the oligonucleotides were annealed by heating at 70°C for 10 minutes followed by cooling to room temperature. The annealed oligonucleotide containing 5' and 3' EcoRI overhanging ends was subcloned into pRS306/HIS3:TetR/Term previously digested with EcoRI. The number

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of copies of inserted oligonucleotide was confirmed by DNA sequencing. The plasmid containing a single copy of the LexA operator was designated pRS306/1xLexAop/HIS3:TetR.

#### D. pRS316/2xLexAop/HIS3:TetR

The subcloning protocol for this construct was the same as described above for pRs306/1xLexAop/HIS3.TetR. The annealed oligonucleotides encoding the LexA operator included overhanging EcoRI ends and during ligation, the individual annealed fragments were able to multimerize, inserting into the parental plasmid more than one copy of the desired LexA sequence. The number of copies of inserted oligonucleotides was confirmed by DNA sequencing.

#### E. pRS306/2xLexAop/HIS3:TetR

A DNA fragment containing two copies of LexA operator and the chimeric HIS3:TetR reporter was excised from pRS316/2xLexAop/HIS3:TetR by digestion with KpnI and BamHI restriction enzymes. The fragment was gel-purified and subcloned into pRS306/Term previously digested with KpnI and BamHI and the resulting construct was sequenced to confirm the presence of two copies of the LexA operator.

#### F. pRS306/4xLexAop/HIS3:TetR and pRS306/8xLexAop/HIS3:TetR

A pair of oligonucleotides SH101A and SH101B were utilized in PCR to amplify the LexA binding site multimer from the plasmid SH18-34ΔSpe [Hollenberg, S.M., et al., Mol. Cell. Biol. 15:3813-3822 (1995)].

SHI01A SEQ ID NO: 20 5'-CCGGAATTCTCGAGACATATCCATATCTAATC

SHI01B SEQ ID NO: 21 5'-CCGGAATTCACTAATCGCATTATCATC

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The amplification product containing four copies of LexA operator was gelpurified, digested with EcoR1, and subcloned into pRS306/HIS3:TetR/Term previously digested with EcoR1. The number of LexA operators were determined by DNA sequencing.

#### G. pRS306/8xLexAop/HIS3::TetR

A PCR strategy was used to link the 5' promoter sequence of the yeast HIS3 gene encompassing nucleotides-75 to +23 to the translational start of TetR. Sequences encoding the SV40 large T antigen nuclear localization signal were fused in frame with the nucleotide sequence encoding the last amino acid residue of TetR. The PCR product was digested with EcoRI and BamHI and inserted into pRS306/Term previously digested with EcoRI and BamHI. The resulting plasmid was designated pRS306/HIS3:TetR/Term, and was shown to encode the complete TetR protein in frame with the nuclear localization signal of SV40 large T antigen. The fusion protein is followed by four amino acids generated by the vector backbone (Arg-Ile-His-Asp).

The LexA binding site multimer from the plasmid pSH18-34\(\Delta\)Spe [Hollenberg, S.M. et al., Mol. Cell. Biol. 15:3813-3822 (1995)] was amplified by PCR, digested with EcoR1, and subcloned into the EcoR1 site of pRS306/HIS3:TetR/Term resulting in plasmid pRS306/8xLexAop/TetR.

#### H. pADH/TetR

The DNA coding sequence of TetR was amplified by PCR from pSLF104 using two oligonucleotides, NcoI-TetR and 3'-TetR terminal primer (SEO ID NO: 17). The resulting product was gel-purified, digested with Ncol and BamHI, and subcloned into a pBTM116 [Bartel, et al., in Cellular Interactions in Development: a Practical Approach, Harrley (ed.), IRL Press; Oxford, pp. 153-179 (1993)] shuttle vector containing an ADH promoter, previously digested with Ncol and BamHI. For construction of this vector, DNA generated by PCR and DNA obtained by restriction enzyme digestion of the polylinker region in plasmid pBluescript (Stratagene, La Jolla, California) were used to engineer additional restriction sites 5' and 3' of the ADH promoter. The TetR protein encoded from this construct is expressed containing additional amino acids Met<sup>2</sup>-Ala<sup>-1</sup> before the initiating methionine and also contains the nuclear localization signal of SV40 large T antigen located after the last amino acid of TetR as described above.

#### pRS306/ADH;TetR/Term

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A fragment encoding the ADH promoter and TetR was removed from plasmid pADH/TetR with Xhol and blunted-ended with the large fragment of DNA polymerase 1 (Gibco BLR, Grand Island, NY). EcoRI linkers (New England BioLabs, Beverly, MA) were added and the fragment was digested with EcoRI and BamHI. The resulting fragment was gel-purified and ligated into pRS306/Term previously digested with EcoRI and BamHI.

#### J. pRS306/4xLexAop/ADH::TetR and pRS306/8xLexAop/ADH::TetR

The subcloning protocol used to insert multiple copies of the LexA operator into pRS306/ADH:TetR/Term was the same as described previously for pRS306/4xLexAop/HIS3:TetR and pRS306/8xLexAop/HIS3:TetR.

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#### III. Plasmids Encoding Binding Proteins

#### A. pLexA-CBD

A DNA fragment containing the CREB binding domain of CBP (CBD), amino acids 461-682, was PCR amplified from plasmid CBP-0.8

[Chrivia, J.C. et al., Nature 365:855-859 (1993)] using a pair of oligonucleotides designated 5' CBD primer and 3' CBD primer.

5' CBD primer SEQ ID NO: 23 5'-GCGAATTCGCCAGGGCAACAGAATGCCACT

3' CBD primer SEQ ID NO: 24 5'-CGGGATCCTGGCTGGTTACCCAGGATGCCTTG

Following gel purification, the amplification product was digested with EcoRI and BamHI, and ligated into plasmid pBTM116 [Bartel, et al., in Cellular Interactions in Development: a Practical Approach, (ed) Hartley, D.A. (IRL Press, Oxford), pp. 153-179 (1993)] previously digested with EcoRI and RamHI

#### B. pVP16-CBD

A DNA fragment encoding the CBP sequence was excised from pLexA-CBD by digestion with EcoRI and BamHI. Plasmid pLexA-CBD was linearized with EcoRI digestion, the resulting overhanging ends blunt-ended using the Klenow fragment of DNA polymerase I, and the ends ligated with BamHI linkers. The resulting fragment was inserted into pVP16 [Hollenberg, et al., Mol. Cell. Biol. 15:3813-3822 (1995)] previously digested with into BamHI.

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#### C. pVP16 CREB

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Plasmid pcDNA3/CREB283 [Sun and Maurer, J. Biol. Chem. 270:7041-7044 (1995)], containing the VP16 transactivation domain fused to sequences of the rat CREB transactivation domain (1 to 283 aa) was linearized with Xhol and BamHI linkers (New England BioLab) ligated to the resulting blunt-ended Xhol sites. DNA encoding the VP16/CREB chimeric protein was removed with HindIII and BamHI digestion and following gel purification, ligated into the HindIII and BamHI sites of pVP16 which encodes the LEU2 gene.

#### D. pVP16-CREB(Bg)II-SacII)-LacZ

A DNA fragment encoding  $\beta$ -galactosidase was PCR amplified from plasmid pSV- $\beta$ -galactosidase vector (Promega, Madison, WI) using a pair of oligonucleotides, 5°  $\beta$ -gal primer and 3°  $\beta$ -gal primer and inserted into the NoiI site of pVP16 to produce pVP16-LacZ.

15 5' β-gal primer SEQ ID NO: 29 5'-ATGGTACCAGCGGCCGCTAGTCGTTTTACAACGTCGTGAC

> 3' β-gal primer SEQ ID NO: 30 5'-ATGGTACCGCGGCCGCTTATTTTTGACACCAGACCAAC

A PCR fragment containing CREB sequences encoding amino acid residues 1 to 283 was amplified from plasmid pRSV-CREB341 [Kwok, et al., Nature 380: 642-646 (1996)] using a pair of oligonucleotides, 5 ° CREB 341 primer and 3 ° CREB 283 primer, and inserted into pVP16-LacZ vector at the BamHI site.

5' CREB 341 primer SEQ ID NO: 25 5'-CGCGGATCCGGATGACCATGGACTCTGGAG

3' CREB 283 primer SEQ ID NO: 28 5'-CGCGGATCCGTGCTGCTTCTTCAGCAGGCTG

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To generate a cassette vector for producing and subcloning mutated CREB sequences as described below, PCR was used to engineer a Bg/II site using oligonucleotides 5' Bg/II primer and 3' Bg/II primer, at nucleotides 273 to 278 and a SacII site using oligonucleotides 5' SacII primer and 3' SacII primer and 3' SacII primer at nucleotides 500 to 505 of the CREB activation domain.

- 5 ' Bg/II primer SEQ ID NO: 31 5 '-CGGAGATCTAAAGAGACTTTTCTCCGGAACTCAG
- 3 · Bg/II primer SEQ ID NO: 32 5 · CGGAGATCTTTACAGGAAGACTGAACTGT
- 5 SacII primer SEQ ID NO: 33 5'-CCACCGCGGCAGTGCCAACCCCGATTTAC
- 3' SacII primer SEQ ID NO: 34 3'-CATCCGCGGTGGTGATGGCAGGGGCTGA

#### E. pLexA-CREB 283

15 A DNA fragment containing the rat CREB transactivation domain (amino acids 1 to 283) was excised from pcDNA/CREB283 [Sun and Maurer, supra] with Smal and Xbal digestion. The 5 'Xbal site was blunt ended with the large fragment of DNA polymerase I (Gibco BRL, Grand Island, NY) and Sall linkers (New England Biolabs, Beverly, MA) added.
20 The fragment was digested with Sall and subcloned into the Sall site of pBTM116.

#### F. plexA-CREB 341

A DNA fragment containing the rat CREB 341 cDNA was amplified by PCR from pcDNA/CREB341 [Kwok, *supra*] using a pair of oligonucleotides, 5 ° CREB 341 primer (SEQ ID NO: 25) and 3 ° CREB 341 primer.

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#### 3 CREB 341 primer SEQ ID NO: 26 5 CGCGGATCCTTAATCTGACTTGTGGCAGTA

After gel purification, the PCR product was digested with BamHI, and subcloned into the BamHI site of pBTM116.

#### G. pLexA-CREB 341-M1

A DNA fragment containing the rat CREB sequence with a mutation changing scrine at position 133 to alanine was amplified by PCR from plasmid Rc/RSV CREB-M1 (Kwok, et al., supra] using the same set of primers as described for pLexA-CREB 341, 5° CREB 341 primer (SEQ ID NO: 25) and 3° CREB 341 primer (SEQ ID NO: 26). The resulting amplification product was gel-purified, digested with BamHI, and subcloned into the BamHI site of pBTM116.

#### H. pVP16-CREB M1

A PCR fragment containing CREB sequences coding for amino acid residues 1 to 283 including the serine 133 mutation to alanine was amplified using a pair of oligonucleotides, 5 ° CREB 283 primer and 3 ° CREB 283 primer (SEQ ID NO: 28). The PCR fragment was gel-purified, digested with BamHI and inserted into the BamHI site of pVP16.

5´ CREB 283 primer SEQ ID NO: 27 5´-CGCGGATCCCCATGACCATGGAATCTGGAGCC

#### pLexA-SRF

A DNA fragment containing human SRF was excised from plasmid pCGN-SRF [Grueneberg, D.A., et al., Science, 257:1089-1095 (1992)] with XhoI and BamHI digestion. The XhoI site of the fragment was blunt-ended by the large fragment of DNA polymerase I (Gibco BRL, Grand Island, NY), ligated with BamHI linkers, digested with BamHI, and inserted

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into pBTM116 previously digested with BamHI.

#### J. pVP16-Tax

A DNA sequence encoding full length Tax protein was excised from pS6424 [Kwok, R.P.S., et al., Nature 380:642-646 (1996)] with BanHI digestion and was inserted into pVP16 previously digested with BamHI.

#### IV. Plasmids For Binding Protein Controls

#### A. pLeu

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Plasmid pVPI6 was digested with *HindIII* and *BamHI* to remove the fragment encoding the VPI6 transactivation domain. The digested vector was blunt-ended and self-ligated.

#### B. pLexA-VP16

The VP16 transactivation domain was PCR amplified from pGal-VP16 [Sadowski, et al., Nature 335:563-564 (1988)] with a pair of oligonucleotides, 5 '-VP16SH and 3 'VP16SH and the resulting amplification product was digested with Clal, blunt-ended, and inserted into pBTM116.

5'-VP16SH SEQ ID NO: 35 GGCTATCGATACGGCCCCCCGACCGAT

3'-VP16SH SEQ ID NO: 36
GCGTATCGATCTACCCACCGTACTCGTC

C. <u>pLexA-Lamin</u>
 See Hollenberg, S.M. et al., Mol. Cell. Biol. 15:3813-3822 (1995)].

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### V. Plasmids Encoding Reporter Gene Controls

#### A. pRS306/Term

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The alcohol dehydrogenase (ADH) terminator sequence was excised from plasmid pBTM116 [Bartel, et al., in Celtular Interactions in Development: a Practical Approach, (ed) Hartley, D.A. (IRL Press, Oxford), pp. 153-179 (1993)] with Sph1 and Pst1 restriction enzymes and both 3'-overhanging sequences were blunted by 74 DNA polymerase (Gibco BLR, Grand Island, NY). The fragment was gel-purified and subcloned into the blunt-ended Notl site in pRS306 [Sikorski and Hieter, Genetics:122:19-27 (1989)]. The orientation of inserted fragment was determined by DNA sequencing.

#### B. pRS316/Term

The subcloning protocol for inserting the ADH terminator sequence into pRS316 was the same as described for inserting the ADH sequence in pRS306.

#### Example 2 Generation of Yeast Assay Transformant

Selection of an appropriate yeast assay strain is an empirical determination based on growth characteristics of the transformed alternatives. A general method to make the appropriate selection is described as follows.

Candidate yeast assay strains were transformed individually with reporter gene constructs and/or a plasmid encoding one of the experimental binding proteins. Assay strains thus transformed were then compared for relative differences in growth characteristics, with an optimal assay strain showing negligible growth on media lacking histidine and vigorous growth on media containing histidine. In practical application of this first step in selection using various plasmids transformed into assay strain Y1584, the following results were observed.

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When the plasmid pLexA-VP16 encoding both the LexA DNA binding domain and the VP16 transactivating domain as a single protein was introduced into the assay cells, growth in the absence of histidine in the media was significantly reduced three days after transformation.

In assays including transformation with plasmids encoding multiple copies of the *tet* operator upstream of the *HIS3* gene, the following plasmids were separately utilized:

pRS303/1xtetop-HIS (encoding a single tet operator sequence), pRS303/2xtetop-HIS (encoding two tet operator sequences), pRS303/3xtetop-HIS (encoding three tet operator sequences), pRS303/4xtetop-HIS (encoding four tet operator sequences), pRS303/8xtetop-HIS (encoding six tet operator sequences), pRS303/8xtetop-HIS (encoding eight tet operator sequences), or pRS303/10xtetop-HIS (encoding tet tet operator sequences).

15 In the assay strains transformed with plasmids encoding either one, two, or three copies of the tet operator upstream from the HIS3 gene, cells grew on media lacking histidine at a rate similar to cells grown on media containing histidine. In yeast assay strains transformed with plasmids encoding either six, eight, or ten copies of the tet operator upstream from the HIS3 gene, cell growth was low suggesting that these strains would not be useful in assays to examine binding and interruption of binding between test proteins. These results suggested that, in assay strains transformed with a reporter plasmid having more than three tet operator sequences upstream from the HIS3 gene, normal activity of the HIS3 promoter is disrupted and that these plasmids

In assays wherein yeast cells were transformed with only reporter plasmids (and not plasmids encoding binding partner fusion proteins) encoding multiple copies of the LexA operator 5° of the TetR gene, the

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following results were observed. Growth of assay cells transformed with plasmids bearing one, two, four, and eight copies of the regulatory LexA operator upstream of the Telt gene appeared to be "copy number" dependent. Yeast cells transformed with plasmids having two copies of the LexA operator grew at a rate significantly higher than those assay cell transformed with a plasmid bearing only one copy of the operator. Cells transformed with plasmids encoding either four or eight LexA operators upstream of the TetR gene grew at an approximately equal rate, and better than assay cells bearing a TetR gene driven by two copies of the operator.

When the alcohol dehydrogenase (ADH) promoter was included upstream of the LexA operator (plasmids encoding either four or eight LexA operators) in the various reporter gene constructs, cell viability was the lowest.

The various cell lines constructed by the methods described above are shown in Table 1, wherein various transformed yeast strains are identified (Strain #) along with the number of LexA operator sequences in the plasmid encoding TetR, the number of tetracycline operator sequences regulating expression of HIS3, and relative growth rate of the transformed strain on media containing histidine. It is important to note that growth variation of transformed cells in media containing histidine is observed, even in cell lines identically transformed. The number of "+" signs in Table 1 is indicative of the host cell's relative ability to grow on media lacking histidine in the absence of transformation with plasmids encoding potential binding proteins. Also in Table 1, a subscript "a" is indicative of transformation with a plasmid bearing the alcohol dehydrogenase promoter; absence of a subscript "a" indicates use of the HIS3 promoter.

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#### Table 1

### Various Yeast Transformants

	Diploids L40	Diploids	1.40
	Strein # LexA TetOp His+		LexA TetOp His+
5	Y1579 1X 2X +++	Y1602	4X, 6X
	Y1581 1X 2X +++	Y1607	4X 6X +++
		Y1628	
	Y1580 2X 2X +++	Y1632	
	YI582 2X 2X +++		
		Y1605	4X, 10X
		Y1610	4X 10X +
10	Diploids L40	Y1622	4X 10X ++
10	Strain # LexA TetOp His+	Y1626	4X_10X ++
	YI583 4X 2X +++		-
	YI585 4X 2X +++	Y1592	8X 2X +++
	YIS87 4X 2X +++	Y1596	8X, 2X +++
	Y1589 4X 2X +++		•
10		Y1598	8X 4X +
15	Y1584 8X 2X +++	Y1635	8X, 4X +
	Y1586 8X 2X +++	Y1637	8X 4X ++
	Y1588 8X 2X +++	Y1601	8X 6X +
	YI590 8X 2X +++	Y1608	8X, 6X +
		Y1629	8X 6X +++
		Y1631	8X 6X +++
	Diploids L40	Y1604	8X 10X +
20	Strain # LexA TetOn His+	Y1611	8X_10X +
	YI591 2X 2X +++	Y1623	8X, 10X ++
	Y1594 2X 2X +++	Y1625	8X 10X ++
	Y1597 2X 4X		on 10n
	Y1633 2X 4X +		
25	Y1636 2X 4X ++	Strain#	LexA TetOp strain His+
		Y1664	4X, 3X w303(50) +++
	Y1600 2X 6X	Y1666	4X, 3X w303(51) +++
	Y1606 2X 6X +		
	Y1630 2X 6X +	Y1668	4X, 2X L40 (69) +++
	Y1627 2X 6X +++	Y1670	4X, 2X L40 (70) +++
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30	Y1603 2X 10X +	Y1665	8X, 3X w303(50) +++
	Y1621 2X 10X ++	Y1667	8X, 3X w303151) +++
	Y1609 2X 10X +	Y1671	8X, 3X L40 (69) +++
	Y1624 2X 10X ++		
		Y1669	8X, 2X L40 (69) +++
35	Y1593 4X, 2X +++	Y1671	8X, 2X L40 (70) +++
33	Y1595 4X 2X +++		
	Y1599 4X, 4X	Y1671	8X <sub>a</sub> 6X L40 (69) +++
	Y1634 4X 4X +		
	Y1638 4X 4X +		
	1 AP 0001		

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#### Example 3 CREB/CBP Binding Interaction

Use of the split-hybrid assay for studies of protein/protein binding wherein one of the binding components is randomly mutagenized was carried out using CREB and CBP binding proteins. The binding of CREB to CBP has been shown to require the phosphorylation of the CREB serine residue at position 133 in a region designated the "kinase-inducible domain" (KID) [Chrivia, et al., Nature 365, 855-859 (1993); Kwok, et al., Nature 370, 223-226 (1994)]. Punctionally, changing serine at position 133 to alanine (a mutant designated CREB-MI) abolishes the ability of CBP to activate CREB-mediated transcription. Preliminary studies have indicated that the CREB-MI mutatu in the split-hybrid system prevents the interaction with CBP and subsequent growth of the yeast assay strain on media lacking histidine. Precisely what other portions of the KID of CREB are required for binding to CBP is unknown, however. To define other potentially important amino acid residues, the KID (amino acid residues 102 to 160) of CREB 341 was randomly mutagenized using PCR.

# A. PCR Mutagenesis and Creation of Mutant Library

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The technique used for mutagenic PCR was a modification of that described by Uppaluri and Towle [Mol. Cell. Biol. 15, 1499-1512 (1995)]. The reaction mixture contained 20 ng of pVP16-CREB(BgIII-SacII)-LacZ, 16 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 67 mM Tris-HCl, pH 8.8, 6.1 mM MgCl<sub>2</sub>, 0.5 mM MnCl<sub>2</sub>, 6.7 μM EDTA, 10 mM β-mercaptoethanol, 1 mM primers, 1mM each dGTP, dTPP, and dCTP, 400 μM dATP, and 2.5 units of Taq DNA polymerase (Promega, Madison, WI). After seven cycles of PCR (94°C for 40 sec, 50°C for 40 sec, and 72°C for 40 sec), the PCR product was amplified a second time using the same primers and Vent DNA polymerase (New England BioLabs, Beverly, MA) under the same conditions for 25 cycles. The resultant PCR product was gel purified, digested with BgIII and

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SacII, and inserted into the BgIII and SacII sites of pVP16-CREB(BgIII-SacII)-LacZ (construction of which is described above). The resulting plasmids were transformed into DH5 $\alpha$  bacterial cells. Transformants were pooled and plasmid DNA was isolated by CsCI gradient centrifugation.

# B. Construction and Use of pVP16-CREB(BgIII-SacII)-LacZ

A DNA fragment encoding the β-galactosidase gene was fused in frame to the carboxyl-terminal end of VP16-CREB as described above. The carboxy-terminal tag allowed identification of clones that contain frameshift and nonsense mutations; colonies that remain positive for  $\beta$ -galactosidase were presumed to contain an open reading frame throughout the mutated region. To facilitate the subcloning of mutated sequences, a cassette version of the CREB cDNA was generated that contained BgIII and a SacII sites flanking the 5' and 3' ends of the KID, respectively. These modifications altered the amino acid residue at position 168 from valine to alanine. The cDNA altered in this manner was indistinguishable from the original VP16-CREB and from VP16-CREB-LacZ when tested in the split hybrid assay. Primers complementary to regions flanking the KID were used in mutagenic PCR amplification reactions as described above under conditions which were optimized to achieve one to three mutations in the 177 bp region encoding the KID. PCR products were introduced into pVP16-CREB(BglII-SacII)-Lac Z in place of wild-type sequence. A library of mutated sequences was transformed into yeast assay strain YI584 expressing LexA-CBD. Approximately 27,000 yeast transformants were screened, yielding about 5,000 colonies that were capable of growing on selective media supplemented with 10 µg/ml of tetracycline and 1 mM of 3AT, determined as described below.

Two screening steps were performed to eliminate uninformative mutations and false positives. First, filter  $\beta$ -galactosidase assays were performed by standard methods [Vojtek, et al., Cell 74:205-214 (1993)] on the 5,000 colonies which exhibited positive growth on media lacking

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tryptophan, histidine, uracil, leucine, and lysine to eliminate expressed proteins having frame-shift and nonsense mutations. Five hundred thirty six colonies developed a dark blue color, whereas 412 colonies turned white and were presumed to express mutants containing either frame-shift or nonsense mutations. The other colonies developed a pale blue color, and control experiments suggested that these colonies may have expressed unstable lacZ fusion proteins. Pate blue colonies were not analyzed further.

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DNA from 536 dark blue colonies was isolated and transformed into E.coli MC1066 cells. One hundred ninety three pVP16-CREB-(BgIII-SacII)-LacZ cDNAs were then isolated.

In a second screening step, the 193 cDNAs were separately retransformed along with pLexA-CBD into the split-hybrid strain as well as into the two-hybrid L40 strain [Vojtek, et al., supra] in order to identify false positives and confirm that the mutant CREB proteins did not interact with CBP. Among the 193 cDNAs re-screened, 152 did not interact with CBP in the yeast two-hybrid system, 15 interacted weakly, and 26 interacted like wild type CREB.

Following these two screening steps, the 152 CREB mutants were sequenced. Seventy CREB mutants were found to contain a single amino acid change. Sixty four CREB mutants contained two amino acid residue mutations and 13 mutants contained more than two amino acid mutations. Mutants containing more than one amino acid alteration were not analyzed further. The expression level of mutant proteins having one amino acid change were determined using a standard  $\beta$ -galactosidase assay.

The CREB mutations identified in the split-hybrid screen were shown to carry amino acid changes centered around the phosphorylation site at serine at position 133. No disrupting mutations were found to contain amino acid alterations outside of the region between amino acids 130 to 141. Most of the mutations abrogated the PKA phosphorylation region, but others were identified at isoleucine position 137, leucine at position 138, and leucine

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at position 141. The mutations at positions 137, 138, and 141 generally changed the hydrophobic residues at these positions to polar residues. The ability of the split-hybrid system to detect only a limited number of CREB mutants, many of which have been proposed previously to disrupt CREB association with CBP [Parker, et al., Mol. Cell. Biol. 16, 694-703. (1996)], indicates the specificity of the split-hybrid system.

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These results lead to interesting suggestions. Various CREB mutations were identified which disrupt CREB-CBP interaction and the majority of disrupting mutations occurred in the CREB PKA phosphorylation motif. This result was consistent with previous observations that unonphosphorylated CREB and CBP do not interact [Kwok, et al., Nature 370:223-226 (1994)]. The most common motif for PKA phosphorylation is an RRX(S/T)X amino acid sequence but RX(S/T)X and KRXX(S/T)X are also phosphorylated [Kemp and Pearson, T.I.B.S. 15, 342-346 (1990)]. The arginine residues in the phosphorylation site are critical for electrostatic interactions with acidic amino acid residues in the catalytic subunit of PKA [Knighton, et al., Science 253, 414-420 (1991)], and consistent with this observation, CREB mutants with changes at arginine residues 130 and 131 were identified in the split hybrid assay that did not interact with CBP.

Results also showed that CREB mutations at amino acids proline at residue 132 and tyrosine 134 were unable to bind CBP. It is likely that the mutations at these residues adversely affect the structure of the phosphorylation motif, although these positions are generally thought to be less critical to CBP binding. It is possible that the substitution of proline at position 132 with threonine created a new phosphorylation site (RXTX) that interfered with the critical phosphorylation of serine at position 133. Although not generally thought to be part of the "classical" consensus PKA phosphorylation motif, hydrophobic amino acids are commonly found carboxy-terminal to PKA sites (Kemp, et al., 7.1.B.S. 19:440-444 (1994)]. The importance of these flanking residues may explain the frequent occurrence

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of disrupting mutations involving tyrosine at position 134. Further studies will be directed to determining if mutations of proline at position 134 and tyrosine at position 134 directly disrupt phosphorylation of serine at position 133 or disrupt binding of CREB to CBP by some other mechanism.

In addition, substitution of serine at position 133 with threonine also prevented the interaction of CREB and CBP. PKA protein substrates containing a phosphorylatable threonine residue are known to exist in nature (i.e., protein phosphatase inhibitor I and myelin basic protein), although they are less common than those with phosphorylatable serines [Zetterqvist, et al., in Peptides and Protein Phosphorylation, (ed.) Kemp, B.E. (CRC Press, Boca Raton, FL), pp. 172-187 (1990)], and synthetic peptides containing serine to threonine substitutions are relatively poor substrates for PKA phosphorylation [Zetterqvist, et al., supra]. In the split-hybrid assay, however, it is unclear whether the mutation of threonine at position 133 disrupts the CREB-CBP interaction or if the mutant fails to become phosphorylated. Despite previous observations that serine residue at position 133 of mammalian CREB can be phosphorylated by a variety of protein kinases other than PKA, for example calcium/calmodulin-dependent protein kinase II and IV, protein kinase C, and a nerve growth factor (NGF)-activated CREB kinase [Sheng, et al., Neuron 4:571-582 (1990); Sheng, et al., Science 252:1427-1430 (1991); Xie and Rothstein, J. Immunol. 154:1717-1723 (1995); Ginty, et al., Cell 77:1-20 (1994)], it is not known which, if any, of these particular protein kinases are able to phosphorylate CREB at the serine at position 133 in yeast. The requirement for integrity of the entire RRXSX amino acid sequence, however, suggests that PKA is a reasonable candidate.

The second category of mutations were identified adjacent the PKA phosphorylation motif. Amino acids isoleucine at position 137 and leucine at position 138 have previously been suggested to be important for hydrophobic interactions of CREB with CBP [Parker, et al., Mol. Cell. Biol. 16. 694-703 (1996)]. In this study, most of the mutations at position 137 and

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138 converted these hydrophobic residues to polar amino acids. Thus, another possibility for the failure of these mutants to bind to CBP is that changes at these positions affect protein folding. Similarly, the mutation at position 141 substituted a polar residue for the wild-type hydrophobic leucine, and this mutation also has the potential to affect protein folding.

Substitution of the isoleucine at position 137 with a hydrophobic phenylalanine residue was found to disrupt the interaction between CREB and CBP as well. This result could have been the result of a detrimental effect on folding because of the steric hindrance associated with the comparatively larger size of phenylalanine. Alternatively, the proposed hydrophobic interactions between CREB and CBP are somewhat specific. Structural studies will be directed to definitively determine how these mutations affect binding.

Perhaps most surprising was the finding that critical mutations

were restricted to a small region in the KID sequence, even though the
relatively low affinity of phosphorylated CREB and CBP, determined to be
between 250 and 400 nM by fluorescence anisotropy measurements [Kwok,
et al., Nature 370, 223-226 (1994)], is consistent with a restricted protein
binding domain. The capability of the split-hybrid system to screen for a

limited number of CREB mutants suggests that the system is highly specific,
and thus, should be useful to identify mutations which disrupt interacts
between other pairs of binding proteins.

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#### Example 4 Tax/SRF Binding Interaction

To further investigate the feasibility of using the split-hybrid system to study protein-protein interactions, a pair of well characterized interacting proteins, SRF and Tax, was tested. Previous studies indicated that SRF and Tax interact in a standard yeast two-hybrid system suggesting that the proteins may be utilized in the split hybrid assay. Plasmid pLexA-SRF, containing a human SRF cDNA fused to the LexA DNA binding domain, was transformed into strain YI584 along with either pVP16-Tax or pVP16 alone. As with the pLexA-VP16 transformation, the yeast strains co-expressing 10 LexA-SRF and VP16-Tax failed to yield any colonies on medium lacking histidine. In contrast, when LexA-SRF was co-transformed with a vector encoding the VP16 activation domain alone, yeast growth occurred on medium lacking histidine, suggesting that TetR expression was not activated. These results demonstrated that a protein-protein interaction in the split-hybrid system can effectively prevent yeast growth and further indicated the utility of the assay for the study of various protein/protein interactions.

#### Example 5 Casein Kinase Binding Assays

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In another example of use of the split hybrid assay to examine protein/protein interactions, Hrr25, a yeast casein kinase isoform, or human casein kinase I isoform  $\delta$ , was employed in the assay with a known binding partner protein

Previous work using the two hybrid assay had identified three genes encoding proteins which interact with the yeast casein kinase isoform Hrr25. Proteins encoded by the genes were designated TIH1, TIH2, and TIH3. The Hri25 expression construct which was generated for use in the two hybrid assay was used in combination with the individual TIH encoding constructs in the split hybrid assay to determine if interaction between the

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binding partners would decrease growth of assay yeast cells on media lacking histidine. Construction of the Hrr25 expression plasmid and isolation of plasmids encoding TIH proteins is discussed below.

In order to identify genes encoding proteins that interact with S. cerevisiae HRR25 CKI protein kinasc, a plasmid library encoding fusions between the yeast GAL4 activation domain and S. cerevisiae genomic fragments ("prey" components) was screened for interaction with a DNA binding domain hybrid that contained the E. coli lexA gene fused to HRR25 ("bait" component). The fusions were constructed in plasmid pBTM116 which contains the yeast TRP1 gene, a  $2\mu$  origin of replication, and a yeast ADHI promoter driving expression of the E. coli lexA protein containing a DNA binding domain (amino acids I to 2021).

Plasmid pBTM116::HRR25 encoding the lex4::HRR25 fusion protein was constructed in several steps. The DNA sequence encoding the initiating methionine and second amino acid of HRR25 was changed to a Smal restriction site by site-directed mutagenesis using a MutaGene mutagenesis kit from BioRad (Richmond, California). The DNA sequence of HRR25 is set out in SEQ ID NO: 39. The oligonucleotide used for the mutagenesis is set forth below, wherein the Smal site is underlined.

# 5'-CCTACTCTTAGG<u>CCCGGG</u>TCTTTTTAATGTATCC-3' (SEO ID NO: 37)

After digestion with Smal, the resulting altered HRR25 gene was ligated into plasmid pBTM116 at the Smal site to create the lexA::HRR25 fusion construct.

Interactions between bait and prey fusion proteins were detected in yeast reporter strain CTY10-5d (genotype=MATa ade2 trp1-901 ku2-3,112 his 3-200 gal4 gal80 URA3::lexA op-lac2.) [Luban, et al., Cell 73:1067-1078 (1993)] carrying a lexA binding site that directs transcription of

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lacZ. Strain CTY10-5d was first transformed with plasmid pBTM116::HRR25 by lithium acetate-mediated transformation [Ito, et al., J. Bacteriol. 153:163-168 (1983)]. The resulting transformants were then transformed with a prev yeast genomic library prepared as GAL4 fusions in the plasmid pGAD [Chien, et al., Proc.Natl.Acad.Sci (USA) 21:9578-9582 (1991)] in order to screen the expressed proteins from the library for interaction with HRR25. A total of 500,000 double transformants were assayed for  $\beta$ -galactosidase expression by replica plating onto nitrocellulose filters, lysing the replicated colonies by quick-freezing the filters in liquid nitrogen, and incubating the lysed colonies with the blue chromogenic substrate 5-bromo-4-chloro-3-indolyl- $\beta$ -D-galactoside (X-gal).  $\beta$ -galactosidase activity was measured using Z buffer (0.06 M Na2HPO4, 0.04 M NaH2PO4, 0.01 M KCI, 0.001 M MgSO<sub>4</sub>, 0.05 M  $\beta$ -mercaptoethanol) containing X-gal at a concentration of 0.002% [Guarente, Meth. Enzymol. 101:181-191 (1983)]. Reactions were terminated by floating the filters on IM Na2CO3 and positive colonies were identified by their dark blue color.

Library fusion plasmids (prey constructs) that conferred blue color to the reporter strain co-dependent upon the presence of the HRR25/DNA binding domain fusion protein partner (bait construct) were identified. The sequence adjacent to the fusion site in each library plasmid was determined by extending DNA sequence from the GAL4 region. The sequencing primer utilized is set forth below.

# 5'-GGAATCACTACAGGGATG-3' (SEQ ID NO: 38)

DNA sequence was obtained using a Sequenase version II kit (US Biochemicals, Cleveland, Ohio) or by automated DNA sequencing with an ABI373A sequencer (Applied Biosystems, Foster City, California).

Four library clones were identified and the proteins they encoded are designated herein as TIH proteins 1 through 4 for Targets Interacting with

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HRR25-like protein kinase isoforms. The TIH1 portion of the TIH1 clone insert corresponds to nucleotides 1528 to 2580 of SEQ ID NO: 40; the TIH2 portion of the TIH2 clone insert corresponds to nucleotides 2611 to 4053 of SEQ ID NO: 41; and the TIH3 portion of the TIH3 clone insert corresponds to nucleotides 248 to 696 of SEQ ID NO: 42. Based on DNA sequence analysis of the TIH genes, it was determined that TIH1 and TIH3 were novel sequences that were not representative of any protein motif present in the GenBank database (July 8, 1993). TIH2 sequences were identified in the database as similar to a yeast open reading frame having no identified function. (GenBank Accession No. 232361, open reading frame YBL0506)

When the various TIH proteins were used in the split hybrid assay in combination with Hrr25, it was observed that Hrr25/TIH3 binding, previously determined to be weaker than Hrr25/TIH2 or Hrr25/TIH1 interactions, produced the lowest level of growth in the transformed yeast

15 strain.

#### CKIδ

In order to isolate cDNAs which encode proteins that interact with CKI\(\delta\), the two hybrid assay was performed using a Lex\(A\)-CKI\(\delta\) fision protein as bait. The coding region of CKI\(\delta\) was subcloned into a \(Bam\)HI site of pBTM116 and transformed into a yeast strain designated CKI\(\delta\)/L40 (MAT a his3 \(\Delta\) 2000 (rpl-901 leu2-3 112 ade2 LYS::(lex\(A\)\)), ltIS3 URA3::(lex\(A\)\) lez GAL 4). CKI\(\delta\)/L4D was subjected to a large scale transformation with a cDNA library made from inouse embryos staged at days 9.5 and 10.5. Approximately 40 million transformants were obtained. Eighty-eight million were plated onto selective media lacking leucine, tryptophan and histidine. The ability of yeast transformants to grow in the absence of histidine suggested that there was an interaction between CKI\(\delta\) and some library protein.

In a second screening, interaction of the two proteins was

assayed by the ability of the interaction to activate transcription of  $\beta$ -galactosidase. Colonies that turned blue in the presence of X-gal were streaked onto media lacking leucine, tryptophan and histidine, grown up in liquid culture and pooled for isolation of total DNA. Isolated DNA was used to transform E. coli strain 600 which lacks the ability to grow on media lacking leucine. Colonies that grew were used for plasmid preparation and three classes of cDNA were identified. One class was closely related to a Drosophila transcription factor dCREBA.

When CKIb/CREB interaction was examined in the split hybrid assay, cells were shown to grow on media containing histidine, but in the absence of histidine, growth was inhibited. Addition of small amounts of tetracycline to the cell culture restored the cell's ability to grow, suggesting that the interaction between CKIô and CREBa was very weak.

#### Example 6 AKAP 79 Binding Assays

Expression Plasmid Utilized

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In still another example of use of the split hybrid assay to examine protein/protein interactions, an anchoring protein for the cAMP dependent protein kinase, AKAP 79, was utilized separately with binding partner proteins including the cAMP protein kinase regulatory subunit type I (RI), the cAMP dependent protein kinase regulatory subunit type II (RII) or calcineurin (CaN). Plasmids used in the assay were constructed as described below.

A 1.3 kb Ncol/BamHI fragment containing the coding region of AKAP 79 was isolated from a pET11d backbone and ligated into plasmid pAS1. Plasmid pAS1 is a 2 micron based plasmid with an ADH promoter linked to the Gal4 DNA binding subunit [amino acids 1-147 as described in Keegan et al., Science, 231:699-704 (1986)], followed by a hemagglutin (HA) tag, polyclonal site and an ADH terminator. The expressed protein was

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therefore a fusion between AKAP 79 and the DNA binding domain of Gal4.

Plasmids encoding RI, RII or CaN were isolated from a pACT murine T cell library in a standard two hybrid assay using the AKAP 79 expression construct described above. Plasmid pACT is a leu2, 2 micron based plasmid containing an ADH promoter and terminator with the Gal4 transcription activation domain II [amino acids 768-881 as described in Ma and Ptashne, Cell, 48:847-853 (1987)], followed by a multiple cloning site. RI, RII and CaN encoding plasmids were isolated as described below.

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A 500 ml SC-Trp yeast cell culture (OD $_{600}=0.6-0.8$ ) was harvested, washed with 100 ml distilled water, and repelleted. The pellet was brought up in 50 ml LiSORB (100 mM lithium acetate, 10 mM Tris pH8, 1 mM EDTA pH8, and 1 M Sorbitol), transferred to a 1 liter flask and shaken at 220 rpm during an incubation of 30 minutes at 30°C. The cells were pelleted, resuspended in 625  $\mu$ l LiSORB, and held on ice while preparing the DNA.

The DNA was prepared for transformation by boiling 400 µ1 10 mg/ml salmon sperm DNA for 10 minutes after which 500 µ1 LiSORB was added and the solution allowed to slowly cool to room temperature. DNA from a Mu T cell library was added (40-50 µg) from a 1 mg/ml stock. The iced yeast cell culture was dispensed into 10 Eppendorf tubes with 120 µl of prepared DNA. The tubes were incubated at 30°C with shaking at 220 RPM. After 30 minutes, 900 µl of 40% PEG<sub>3350</sub> in 100 mM Li acetate, 10 mM Tris, pH 8, and 1 mM EDTA, pH 8, was mixed with each culture and incubation continued for an additional 30 minutes. The samples were pooled and a small aliquot (5 µl) was removed to test for transformation efficiency and plated on SC-Leu-Trp-His media and grown for one hour at 30°C with shaking at 220 RPMS. Harvested cells were resuspended in 5.5 ml SC-Leu-Trp-His containing 50 mM 3AT (3-amino triazole) media and 300 µl aliquots plated on 150 mm SC-Leu-Trp-His also containing 50 mM 3AT. Cell were left to

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grow for one week at 30°C.

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After four days, titer plates were counted and  $1.1 \times 10^5$  colonies were screened. Large scale  $\beta$ -gal assays were performed on library plates and ten positive clones were isolated for single colonies. One of these colonies grew substantially larger than the rest, and was termed clone 11.1. Sequence from clone 11.1 revealed an open reading frame 487 aa long which was correctly fused to the Gal-4 activation domain of pACT. The NIH sequence database was searched and the sequence was found to be closely homologous to the human calmodulin dependent protein phosphatase, calcineurin.

Additional screening using pACT Mu T-cell library DNA and the pASI AKAP 79 bait strain was performed in order to identify other AKAP 79 binding proteins by the protocol described above. Results from screening approximately 211,000 colonies gave one positive clone designated pACT 2-1. Sequencing and a subsequent data base search indicated that the clone had 91% identity with rat type  $1\alpha$  regulatory subunit of protein kinase A (RI).

The library was rescreened using the same AKAP 79 bait and fifteen positives were detected from approximately 520,000 transformants. Of these fifteen, eleven were found to be homologous to the rat regulatory subunit type I of PKA. Each of these isolates were fused to the 5' untranslated region of RI and remained open through the initiating methionine.

#### Split Hybrid Analysis

In split hybrid analysis of AKAP79 binding interactions, a plasmid was first constructed for expression of a LexA:AKAP 79 fusion protein. An AKAP 79 coding region was excised from pAS AKAP 79 as an NcollBamHI fragment and inserted into pBTM116 previously digested with the same enzymes. The resulting plasmid was designated pBTM116-AKAP79.

Approximately 50,000 W303 yeast cells (strain YI665, see Table 1) in logarithmic growth were rinsed in media lacking histidine, suspended in 100  $\mu$ l to 200  $\mu$ l of the same media, and plated on agar lacking

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histidine (to select for absence of protein/protein interaction) and also lacking leucine and tryptophan (to select for transformants bearing expression constructs encoding AKAP 79 and its binding partner). When RII was employed as the AKAP 79 binding partner, 2 to 4  $\mu$ M tetracycline and 5 mM 3AT were required to prevent the transformed host from growing under conditions where the expressed proteins interacted.

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Once conditions were established under which growth of the transformed host was eliminated, various candidate inhibitor compounds were separately added to the agar. It was presumed that if one of the candidate compounds was capable of disrupting AKAP 79 interaction with the binding partner protein, growth of the transformed host should be detectable in the vicinity of the compound on the agar. In the split hybrid assay wherein AKAP 79 and RII binding was examined,  $2\mu$ I of a 30 mM stock solution of ICOS Compound 4273 in DMSO,  $2\mu$ I of a 10 mM stock solution of ICOS Compound 1062 in DMSO, and  $2\mu$ I DMSO alone (as a negative control) were spotted on to the plate which was incubated at 30°C for four to five days. For ICOS Compound 4273 a ring of growth was detected.

In order to determine an  ${\rm IC}_{50}$  for an inhibitor identified as described above, alternative methods may be used. In one method, the inhibitor compound is added to the agar over a range of concentrations. Ideally, the compound is diluted to the point that host cell growth is essentially not detectable.

In another method, a 96 well plate is used and the compounds of interest are serially diluted across one row of a 96 well plate, one compound per row. Media lacking histidine, tryptophan, and leucine is added (presuming that the expression plasmids encoding the binding partners also encode trp and leu proteins) along with the appropriately transformed host yeast strain. Tetracycline and 3AT are added at concentration previously determined to extinguish growth of the transformed host cell. After two to five days incubation at 30°C, the plate wells are read at approximately 600

nm using a plate reader. The concentration of inhibitor half way between zero and the lowest concentration that permits growth of the host cell to the level observed on media containing histidine is estimated to be  $IC_{50}$ .

A modification of this second method is particularly amenable

5 for use in a high throughput screen of large numbers of candidate inhibitors.

For example, rather than attempting to determine the IC<sub>50</sub> for a previously identified inhibitor, separate candidate inhibitors are added to each well of a 96 well plate, preferably at more than one concentration, and host cell growth determined after several days incubation. Inhibitory activity of compounds identified in this manner is confirmed on an agar plate and the IC<sub>50</sub> determined on 96 well plates, each assay as described above.

# Example 7 General Application of The Split-Hybrid Screen

In order to examine general utility of the split hybrid system,

various experiments were conducted with binding proteins known to interact.

In addition, a number of control experiments were included in order to determine if the effects observed with the known binding partners were in fact due to protein/protein interaction.

## A. Yeast Assay Strain Construction

Yeast transformants used in assays indicated below were derived from LYS2-deficient strains AMR69 (Mat a his3 lys2 leu2 trp1, URA3:LexA::LacZ) and AMR70 (Mat α his3 lys2 trp1 leu2, URA3:LexA::LacZ) [Hollenberg, et al., Mol. Cell. Biol. 15, 3813-3822 (1995); Chien, et al., Proc. Natl. Acad. Sci. (USA) 88:97578-9582 (1991); Fields and Song, Nature 340:245-246 (1989)]. Yeast were grown in YEPD or selective minimal medium using standard conditions [Sherman, F., et al., Mcthods in Yeast Genetics, Cold Spring Harbor Lab., Cold Spring Harbor, NY (1986); Methods in Enzymology, Vol. 194 Guide to Yeast Genetics and

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Molecular Biology. Eds. Christine and Fink]. Derivatives of both AMR69 and AMR70 strains lacking URA3 were first generated by streaking cells on synthetic media containing 5 mg/ml 5-fluoro-orotic acid (5FOA) [Methods in Enzymology, Vol. 194 Guide to Yeast Genetics and Molecular Biology. Eds. Christine and Fink]. Two URA3 deficient mutants were required due to the fact that these strains were subsequently mated. URA3-deficient colonies were confirmed by testing for uracil auxotrophy and deletion of the URA:LexA::LacZ locus was confirmed by an absence of  $\beta$ -galactosidase activity assayed by standard methods. The mutant strains selected were designated 69-4 and 70-1.

Targeted integration of pRS306/8xLexAop/TetR was carried out by transforming [Hollenberg, et al., Mol. Cell. Biol. 15, 3813-3822 (1995)] the 69-4 strain with plasmid linearized at a unique Ncol site. The reporter gene construct was constructed using parental plasmid pRS306 which encodes URA3 as a selectable marker. Stably integrated plasmid thereby permitted selection on media lacking uracil. The positive uracil prototrophic strains were examined by Southern analysis to confirm insertion of the plasmid sequences.

Targeted integration of pRS303/2xtetop-LYS was carried out by transformation [Hollenberg, et al., supra] of strain 70-1 with plasmid linearized at a unique Hpal site. The resulting lysine prototrophic strains were examined by Southern analysis to confirm insertion of the plasmid DNA.

The AMR69 derivative strain (MAT  $\alpha$ ) containing the pRS303/2xetop-LYS insertion was mated with the AMR70-derivative strain (MAT  $\alpha$ ) containing pRS306/8xLexAop/TetR and mated cells were selected on media lacking both lysine and uracil. Single colonies were grown up and tested for the ability to grow on media lacking histidine. The resulting strain was designated YI584. In instances where yeast strains were transformed with other reporter gene pair combinations, the strains were uniquely designated.

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subsequently transformed [Hollenberg, et al., supra] with plasmids encoding chimeric binding protein. Plasmids encoding the LexA DNA binding region were generally derived from parental plasmid pBTM116 which also encodes TRPI as a selectable marker. Plasmids encoding the VP16 transactivating domain were generally derived from parental plasmid pVP16 which also encodes LEU2 as a selectable marker. Yeast cells which were successfully transformed with the four exogenous plasmids were therefore selected by an ability to grow on media lacking lysine, uracil, tryptophan, and leucine. Plasmids encoding various binding proteins were transformed into the yeast assay strain as indicated below.

#### B. Liquid Assay

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After three days growth at 30°C on selection media as described above, a pool of colonies from each transformation was collected and diluted in 5 ml selective media. The mixture was vortexed and immediately sonicated for ten seconds. Cells in the resulting suspension were counted and seeded at 1000 cells/ml in selective media, 2 ml per 15 ml tube. Tetracycline, 3AT, and histidine were included as determined appropriate by the method described above. Each aliquot of cells was incubated with shaking for two days at 30°C and cell density measured at ODcoo.

## 20 C. Characterization of the Assay

The utility of the split-hybrid assay was first determined using well characterized binding proteins and various controls.

In an initial study, YI584 cells were transformed with plasmids pLexA-VP16 and pLeu. While the expressed proteins from the two plasmids do not interact, pLexA-VP16 encodes a fusion protein containing the VP16 activation domain fused directly to LexA which contains a DNA binding domain. The chimeric LexA-VP16 protein is a strong transactivator for a promoter containing LexA operators. Plasmid pLeu is essentially a blank used

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## as a control co-transformation plasmid.

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Yeast transformed with the LexA-VP16 plasmid were able to express TetR protein as indicated by gel shift analysis using a tet operator oligonucleotide. In addition, the cells were unable to grow on media in the absence of histidine. Combined, these observations suggested that overexpressed TetR protein was capable of binding to tet operators and preventing the expression of HIS3. The transformed yeast grew on plates containing histidine, further indicating that overexpression of TetR did not have a toxic effect on the assay cells.

The results were consistent with previous observations and supported the earlier suggestion that activation of TetR expression, either through a single transcription factor or association of individual transcription factor domains, is capable of preventing assay cell growth on media lacking histidine, presumably by eliminating HIS3 production.

Example 8

Split-Hybrid Assay With Weakly Interacting Binding Proteins

Protein/protein interaction was examined in the split-hybrid assay to determine utility of the system using two fusion proteins known to

interact weakly. In this instance, the binding proteins were a 283 amino acid fragment of a cAMP regulatory binding protein (CREB283) fused to LexA and a fragment of the CREB binding protein consisting of the CREB binding domain (CBD) fused to VP[6.

In this assay, yeast strain Y1584 described above was employed and transformation carried out as previously described. In a first assay, plasmids pLexA-CREB and pVP16-CBD were transformed into the cells and cell growth was observed in the absence of histidine in the media. Expression of the fusion proteins was confirmed by Western blotting. Attempts to decrease cell growth by titration with 3AT were unsuccessful in that the concentration of 3AT required to reduce growth in cells transformed with

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pLexA-CREB and pVP16-CBD also eliminated growth in cells transformed with pLexA-CREB and the control plasmid pVP16.

In light of these results, two alternative approaches were taken in order to permit study of binding proteins wherein the interaction is relatively weak. Under the assumption that the system was failing at the level of TetR transcription, alternative approaches were taken in attempts to amplify the TetR effect on expression of HIS3 gene. To achieve this end, assay cells were transformed with reporter constructs which encoded multiple tet operator sequences upstream from the HIS3 gene. In the second approach, the HIS3 promoter used to drive expression of the TetR gene was replaced with the stronger alcohol dehydrogenase (ADH) promoter.

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In YI596 cells wherein the ADH promoter replaced the HIS3 promoter to drive TetR expression, transformation with plasmids pLexA-CREB and pVP16-CBD showed substantially decreased growth on his media as compared to that in assay strain YI592 wherein the HIS3 promoter was used to drive TetR expression. However, in cells transformed with plasmids pLexA-CREB 341-M1 and pVP16-CBD, no decrease in assay cell growth was detected on media lacking histidine. These results indicate that incorporation of the ADH promoter to drive TetR expression may be more useful in studies involving binding proteins that have low affinity.

When assay strains were utilized which incorporated plasmids wherein expression of the HIS3 gene was driven by multiple copies of the tet operator, transformed cell lines did not grow well enough to indicate potential utility in subsequent assays.

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#### Example 9 General Assay Methods

#### A. "Fine Tuning"

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In instances where either of the test fusion proteins possesses intrinsic capacity for transcriptional activation, TetR will be expressed and growth of the assay strain media lacking histidine will be depressed proportional to the level of TetR expression. In order to restore growth of these cells to approximately the level observed on media containing histidine, the initially transformed assay yeast strains are grown in the presence of increasing concentrations of tetracycline which binds to the TetR gene product and prevents TetR binding to the tet operator. Precise titration of expressed TetR with tetracycline, only to the point that growth of the assay strain is restored to the level detected in the presence of histidine, permits detection of subsequent decreased growth of the assay strain following increased TetR expression resulting from interaction of the test binding proteins. The empirically determined tetracycline concentration is therefore employed to increase "signal-to-noise" ratios under assay conditions.

After an appropriate tetracycline concentration has been determined for each of the candidate assay strains, the cells are transformed with the second plasmid encoding the second fusion binding protein. As before, growth of each candidate assay strain is examined on media in the presence and absence of histidine. A desirable yeast assay strain is chosen which shows vigorous growth in the presence of histidine and negligible growth on media lacking histidine (indicative of the expected protein/protein interaction and resultant decreased expression of HIS3).

In instances where binding between the two test proteins is comparatively weak. TetR expression may not be sufficiently increased to abolish HIS3 expression and cells expressing the resultant low levels of HIS3 will still grow on media which lacks histidine. Cells which show this low level of viability are grown in the presence of increasing concentrations of 3-

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aminotriazole (3AT), a competitive inhibitor in the histidine synthesis pathway, in order to reduce cell growth to negligible levels when plated on media lacking histidine. As with titration of TetR with tetracycline, addition of 3AT to the media is designed to increase the signal-to-noise ratio by providing significant changes in growth in the presence and absence of histidine in the media.

In a practical application of the methods for fine tuning, binding between CREB and the CREB binding protein (CBP) is illustrative. Growth of the yeast strain Y1584 transformed with pLexA-CBD, encoding the CREB binding domain (CBD) of CBP, and pVP16-CREB or pLexA-CBD and the control plasmid pVP16 was substantially decreased and virtually indistinguishable growth rates were detected in both instances on media lacking histidine. This observation indicated that the LexA-CBD protein product possessed sufficient transactivating capacity to eliminate HIS3 production. In order to distinguish growth differences between assay cells transformed with either pVP16 and pVP16-CREB, increasing amounts of tetracycline were added to the media lacking histidine.

In both transformants, tetracycline was able to relieve growth repression in a dose dependent manner, and at increasing concentrations of tetracycline, the difference in growth between the two colonies was increasingly magnified, with the most distinct growth difference observed following addition of tetracycline at 10  $\mu$ g/ml. Addition of tetracycline was therefore able to overcome the intrinsic transactivating capability of the LexA-CBD fusion protein.

Because the ultimate use of the split-hybrid system is for structure-function studies, mutagenesis studies, drug identification and library screens, it is important to minimize background growth that might be confused with disrupted protein-protein associations. This can be accomplished by the addition of 3AT, a competitive inhibitor of the HIS3 gene product. For instance, in the presence of 10 µg/ml of tetracycline, the yeast strain

transformed with pLexA-CBD and pVP16-CREB still conferred approximately 12% growth of that observed in the presence of his + media. To diminish this background, increasing concentrations of 3AT were added to the media in the presence of 10 µg/ml of tetracycline. At the 3AT concentration of 0.25 mM, the growth of the yeast strain expressing LexA-CBD and VP16-CREB was below 5%, while the growth of the control strain was still maintained at 70% of control levels. These results indicate that split-hybrid system can be modulated by 3AT in addition to tetracycline in order to effectively increase the signal-to-noise ratio.

#### B. Preparation of yeast extracts

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In order to assess the utility of various plasmids to function in the split-hybrid assay, a number of control experiments can be employed which lend insight into expression of a desired protein from the transformed plasmid. For example, standard immunological methodologies, i.e., immunoprecipitation, ELISA, etc., can be used to determine to the extent to which a desired protein is expressed. Similarly, a variation of the gel shift assay (discussed immediately hereafter) can be used to determine both if a protein is expressed and if the expressed protein is capable of DNA binding. In each of these control assays, a yeast extract is required which can be prepared as follows.

Extracts were prepared as described by Uppaluri and Towle [Mol. Cell. Biol. 15:1499-1512 (1995)] and were used for electropheretic mobility shift assays as discussed below. The yeast cells transformed with pLexA-VP16 were grown in 100 ml of selective synthetic medium lacking uracil, tryptophan, and lysine to a density of  $A_{600} = 1$ . Cells were harvested and washed with 5 ml of EB (containing 0.2 M Tris-HCl, pH 8.0, 400 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 10 mM MgCl<sub>2</sub>. 1 mM EDTA, 10% glycerol, and 7 mM  $\beta$ -mercaptoethanol). Cells were transferred to microcentrifuge tubes and collected by centrifugation. After resuspending in 200  $\mu$ l EB containing 1

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mM phenylmethylsulfonyl fluoride (PMSF),  $1\mu g/ml$  leupeptin, and  $1\mu g/ml$  pepstatin, a one-half volume of glass beads was added. The suspension was frozen in a -80°C freezer for 1 hour and thawed on ice. Thawed cells were vortexed at 4°C for 20 minutes, after which an additional  $100~\mu l$  EB was added, and cells were left on ice for 30 minutes. The suspension was centrifuged for 5 minutes, the supernatant was transferred to a new tube which was centrifuged for 1 hour in a microcentrifuge. The supernatant was then made to 40% with  $(NH_4)_2SO_4$  and gently rocked for 30 minutes. After a 10 minute centrifugation, the pellet was resuspended in  $300~\mu l$  of 10~mM HEPES, pH 8.0, 5 mM EDTA, 7 mM  $\beta$ -mercaptoethanol, 1 mM PMSF, 1  $\mu g/ml$  leupeptin, and 1  $\mu g/ml$  pepstatin, and 20% glycerol. The resulting suspension was dialyzed against the same buffer, and aliquots were stored at  $80^{\circ}C$ .

# Electrophoretic mobility shift assays

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Electrophoretic mobility shift assays were performed as described by Shih and Towle [J. Biol. Chem. 267:13222-13228 (1992)]. Double-stranded tet operator oligonucleotides were prepared by combining equivalent amounts of complementary single-stranded DNA (SEQ ID NOS: 7 and 8) in a solution containing 50 mM Tris-HCl, pH 8.0, 10 mM MgCl<sub>2</sub>, and 50 mM NaCl<sub>2</sub>, heating the mixture to 70°C for 10 minutes, and then cooling to room temperature. The annealed oligonucleotides were labeled by filling in overhanging 5° ends using the Klenow fragment of E. coli DNA polymerase I with [α-3²P]dCTP. Binding reactions were carried out in 20 μl containing 10 mM Tris-HCl, pH 7.5, 50 mM NaCl, 1 mM EDTA, 1 mM dithiothreitol, 5% glycerol, and 2 mg of poly[d(I·C)]. A typical reaction contained 20,000 cpm (0.5-1 ng) of end-labeled DNA with 3-5 μg of yeast extract. Following incubation at 22°C for 30 minutes, samples were separated on a 4.5% nondenaturing polyacrylamide gel containing 50 mM Tris, 384 mM glycine, and 2 mM EDTA, pH 8.3. For competition binding experiments, the

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conditions were exactly as above except that specific and nonspecific competitor DNAs were included in the binding mixture before the yeast extract was added. The concentration of tetracycline, a competitive inhibitor of TetR/tet operator binding, was 1  $\mu$ M when utilized.

# Example 10 Application of the Split-Hybrid Assay to Identify Agents That Prevent Receptor Desensitization and Drug Tachyphylaxis

Over half of the drugs that are used clinically affect the function of seven transmembrane receptors. Although many of the characteristics of these receptors are distinct, two general features appear to be conserved. One is the ability to signal through dissociation of heterotrimeric G proteins. The second is the capacity to lose responsiveness to ligand binding in a process termed desensitization which is mediated by receptor phosphorylation and the subsequent binding of factors that recognize the phosphorylated state of the receptor which prevents continued signaling. Desensitization results in an intrinsic limitation to drug action imposed by the action of the drug itself, i.e., activation of a receptor by a hormone or drug initiates mechanisms that prevent subsequent responses to repeated administration of the same agent. The coupled mechanisms of activation and deactivation together have been termed "homologous desensitization," while the inability of a drug to maintain its efficacy is known as "tachyphylaxis." Even though the mechanisms underlying homologous desensitization have been worked out in great detail over the past few years, there are currently no useful pharmacological approaches available that prevent the inactivation mechanism.

The potential clinical utility of agents that could prevent or modulate drug desensitization is enormous. Four examples where therapy is limited by the inability of receptors to maintain responsiveness to drugs include: (i) asthma wherein desensitization of airway adrenergic receptors

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renders epinephrine treatment ineffective after a period of hours; (ii) congestive heart failure wherein desensitization of adrenergic and VIP receptors, coupled with an elevation of the  $\beta$  adrenergic receptor kinase ( $\beta$ ARK), prevents the inotropic effects of endogenous regulatory hormones; (iii) Parkinson's disease, wherein dopamine receptor desensitization limits the usefulness of agents like L-Dopa; and (iv) chronic pain wherein tolerance results from opiate receptor desensitization. Indeed, it is difficult to conceive of a pharmacological modality in use today that is not limited in its effectiveness by the phenomenon of desensitization.

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The biochemical basis for G protein-coupled receptor desensitization involves three classes of proteins including arrestins, kinases and Gproteins, all of which have been cloned [Lefkowitz, Nature Biotechnology 14:283-286 (1996)]. Following activation of a seven transmembrane receptor, a region is phosphorylated by one or more G protein-coupled receptor kinases (known as GRKs 1-6). For example, in the  $\beta$ -adrenergic receptor ( $\beta$ AR) and rhodopsin, the cytoplasmic tail is phosphorylated [Premont, et al., J. biol. Chem. 269:6832-6841 (1994): Freedman. et al., J. Biol. Chem. 270:17953-17961 (1995); Palczewski, et al., J. Biol. Chem. 266:12949-12955 (1991); Palczewski, et al., J. Biol. Chem. 270:15294-15298 (1995)] while in the m2 muscarinic receptor, the third cytoplasmic loop is phosphorylated [Nakata, et al., Eur. J. Biochem. 220:29-36 (1994)]. The best characterized members of the family of G protein receptor kinases are the  $\beta$ AR kinase ( $\beta$ ARK) and rhodopsin kinase which are both membrane-associated. While rhodopsin kinase contains an intrinsic membrane targeting signal [Inglese, et al., Nature 359:147-150 (1992)], \(\beta\)ARK appears to be targeted to the membrane by association with G protein  $\beta_Y$  subunits [Pitcher, et al., Science 257:1264-1267 (1992); Inglese, et al., Nature 359:147-150 (1992)]. Once the substrate receptor for each kinase is activated, presumably by ligand binding, the kinase associates and phosphorylates serine and threonine residues on the receptor. The phosphorylated receptor then becomes a binding target for one or more

other proteins. In the case of  $\beta$ AR, for example, phosphorylation allows binding of arresting which prevents association with G proteins and promotes receptor sequestration and desensitization. Using the  $\beta$ AR as an exemplary desensitization model, it becomes apparent that multiple steps in the pathway appear to provide potential points of regulation each of which is amenable to the split-hybrid screen to identify molecules that can block the overall desensitization pathway. Specifically in the case of  $\beta$ AR, the split hybrid system can be used to identify small molecules that: (i) prevent interaction between  $\beta$ ARK and the G protein  $\beta$  subunit; (ii) inhibit  $\beta$ ARK activity; and (iii) disrupt the  $\beta$ ARK:arresting complex.

#### A. Plasmid Constructions

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The study of G-protein receptor kinases in the split-hybrid system involves three or more recombinant proteins or two or more recombinant proteins and a recombinant peptide library. In the split-hybrid system discussed above, two yeast primary expression plasmids are employed: pBTM116 [Bartel et al., Cellular Interactions in Development: a Practical Approach. (cd) Hartley, IRL Press, Oxford, pp. 153-179 (1993)], which encodes the LexA-fusion protein and the TRPI selectable marker, and pVP16 [Hollenberg et al., Mol. Cell. Biol., 15:3813-3822 (1995)], which encodes the VP16-fusion protein and the LEU2 selectable marker. In order to study interactions involving more than two recombinant proteins in the split-hybrid system, however, additional selectable markers are employed. Construction of additional yeast expression plasmids which are used to examine interactions between more than two binding proteins is discussed below.

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#### Plasmid pDRM

A DNA fragment comprising the ADH promoter and LexA sites, the TetR encoding gene, the nuclear localization signal, and the ADH tenninator sequence are removed from pRS306/4xLexAop/ADH::TetR with SacI. blunt-ended, and digested with SalI. The fragment is isolated and ligated into pRS303/2xtetop-LYS2 which has previously been digested with NoII, blunt-ended, and digested with SalI. The resulting plasmid, designated pDRM, is integrated into the LYS2 locus in the yeast genome as described above, and the resulting strain designated YIDRM. Placing the repressor gene and selectable marker reporter gene in the LYS2 locus allows ER43 to be used a selectable marker.

#### 2. Plasmid pRSURA3

A modified version of the pRS306 vector [Sikorski et al., Genetics, 122:19-27 (1989)] containing the URA3 selectable marker gene is also used to encode additional recombinant proteins in the split-hybrid system. The plasmid, pRS426, has the 2 micron origin of replication inserted into a unique AaII site of pRS306. Plasmid pRS426 is further modified in the following manner:

- (i) The ADH promoter sequence is amplified by PCR from BTM116 using primers which incorporate into the amplification product the DNA sequence encoding the SV40 large T antigen nuclear localization signal (NLS) and an initiating ATG sequence 3' to the ADH promoter. The ADH promoter/NLS/ATG sequence is inserted into the polylinker of pRS426.
- (ii) The ADH terminator sequence is amplified by PCR from 25 BTM116 using primers which incorporate into the product a DNA sequence encoding an antibody tag, for example, FLAG, hemagglutinin protein (HA), or thioredoxin (Thio) (FLAG, HA, and Thio antibodies are available through Santa Cruz Biotechnology, Santa Cruz, CA) and DNA sequences encoding stop codons in all three frames to the 5' end of the ADH terminator sequence.

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The antibody tag/stop codon/ADH terminator sequence is inserted into the polylinker of pRS426.

#### 3. Plasmid pRSADE2

PCR is used to engineer unique restriction sites, including for example, Bg/III, Eco47III, MluI, Nhel, and SphI, immediately adjacent the 5' and 3' ends of the URA3 cassette in pRSURA3. The URA3 cassette is digested from pRSURA3 and replaced with the ADE2 cassette which is amplified by PCR.

#### 4. Plasmid pBTM116/AD4

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A fragment containing the ADH promoter, polylinker, and ADH terminator is digested from pAD4 [Young et al., Proc. Nat'l. Acad. Sci. (USA), 86:7989-7993 (1989)] with BamHI, blunt-ended and inserted into the blunt-ended Pvul site of BTM116 as described [Keegan et al., Oncogene, 12:1537-1544 (1996)], and the resulting vector designated pBTM116/AD4. PCR is also used to engineer a nuclear localization signal 3' of the ADH promoter as described above. This vector contains the TRP1 selectable marker and can encode two recombinant proteins: (i) a LexA-fusion protein and (ii) a protein expressed from the pAD4 region of the vector.

# B. βARK and G Protein β Subunit Binding

In a first application of the split hybrid assay, disruption of binding between the carboxy-terminal domain of  $\beta$ ARK, containing the pleckstrin homology (PH) domain, and the G protein  $\beta$  subunit  $(G\beta_2)$  is examined. Previous work indicates that the PH domain of  $\beta$ ARK interacts directly with the  $\beta\gamma$  subunits of G proteins [Pitcher, J.A., et al. Science 257:1264-1267 (1992) and Touhara, K. et al., J. Biol. Chem. 269:10217-10220 (1994)]. Consistent with this observation is work by Pumiglia, et al. [Pumiglia, K.M., et al., J. Biol. Chem. 270:14251-14254 (1995)] which

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indicates that  $G\beta_2$  interacts with Raf1 in yeast and that the interaction is disrupted by  $\beta$ ARK in vitro.

A DNA fragment containing the carboxy-terminal 222 amino acids (residues 467 to 689) of  $\beta$ ARK1, which includes the PH domain, is amplified by PCR from bovine  $\beta$ ARK1 [Pitcher et al., Science, 257:1264-1267 (1992)] and the gel-purified amplification product is inserted into pBTM116. The resulting plasmid is designated LexA-COOH- $\beta$ ARK. A DNA fragment containing the entire coding sequence of  $G\beta_2$  [Fong et al., Proc. Nat'l. Acad. Sci. (USA), 84:3792-3796 (1987)] is PCR amplified from pGEM-112I( $\gamma$ -)G $\beta_2$  [Inigez-Lluhi et al., JBC, 267:23409-23417 (1992)] and the gel-purified amplification product inserted into pVP16. The resulting plasmid is designated pVP16-G $\beta_2$ . PCR is used in a similar manner to clone the carboxy-terminal domain of  $\beta$ ARK into pVP16 and G $\beta_2$  into pBTM116.

 $\beta$ ARK and  $G\beta_2$  binding is first examined in the two-hybrid system to determine if expression of either binding partner as a fusion protein in yeast affects protein/protein interaction. Binding of the two proteins is then examined in the split hybrid assay in order to determine if protein/protein interaction is capable of abolishing growth of the assay yeast strain. As above, addition of tetracycline and/or 3-aminotriazole required to maximize the difference in growth in the presence and absence of histidine is empirically determined

Split-hybrid yeast strains containing  $\beta$ ARK and  $G\beta_2$  subunits are used to screen libraries of small molecules. Several types of small molecule libraries can be examined in the split-hybrid assay, including for example, chemical libraries, libraries of products naturally produced by microorganisms, animals, plants and/or marine organisms, combinatorial, recombinatorial, peptidomimetic, multiparallel synthetic collection, protein, peptide and polypeptide libraries. A library of small peptides can be cloned into pRSURA3 as described [Yang et al., Nuc. Acids Res. 23:1152-1156 (1995) and Colas et al., Nature, 380:548-550)]. Peptides corresponding to the

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carboxy-terminus of  $\beta$ ARK or other GRKs which have previously been shown to block calcium channel desensitization in intact neurons, presumably by blocking  $\beta$ ARK and  $G\beta_2$  binding and subsequent trafficking of  $\beta$ ARK to the cellular membrane [Diverse-Pierluissi, et al., Neuron 16:579-585 (1996)] can be identified in such a screen. Further, it is important to show that the molecules identified through the split hybrid selection affect  $\beta$ ARK: $G\beta$  interaction as opposed to, for example, tetracycline analogues identified in the screen that would not be useful to specifically modulate  $\beta$ ARK: $G\beta_2$  binding.

# B. Identification of βARK Inhibitors

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In a second approach, agents that directly inhibit \$\textit{\textit{BARK}}\$ function are identified in a modification of the split-hybrid system. White identification of specific \$\textit{\textit{BARK}}\$ inhibitors may be difficult, preliminary data from split hybrid assays using CREB/CBP binding partners indicates that the system can be used to identify serine kinase inhibitors. The serine kinase results also suggest several approaches can be employed in attempts to overcome potential problems in identifying \$\textit{\textit{BARK}}\$ inhibitors.

Briefly, binding between the phosphorylated G-protein coupled receptor (P-GR) and arresting is examined first in the standard two hybrid assay, followed by identification of inhibitors of P-GR/arresting binding in the split hybrid assay. For these studies, fragments of three G protein-coupled receptors are examined: the carboxy-terminal tail of  $\beta_2$ AR and the third cytoplasmic loop of the m2 muscarinic receptor. A DNA fragment containing the carboxy-terminal tail of the  $\beta_2$ AR (amino acids 330 to 413) is PCR amplified (Kolbilka et al., JBC, 262:7321-7327 (1987)] and the gel purified product inserted into pBTM116/Ad4 to produce a LexA- $\beta_2$ AR fusion gene. The resulting plasmid is designated pBTM- $\beta_2$ AR/AD4. A DNA fragment containing the third cytoplasmic loop of the human m2 muscarinic receptor (nucleotides 268-324) is amplified from pGEX-13m2 [Haga et al., JBC,

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269:12594-12599 (1994)] by PCR and cloned into pBTM116/Ad4 creating a LexA-m2 fusion gene. The resulting plasmid is designated pBTM-m2/AD4. The entire bovine βARK1 coding sequence [Benovic et al., Science, 246:235-240 (1989)] is PCR amplified and cloned into the polylinker region originating from AD4 in pBTM-β2AR/AD4 and pBTM-m2/AD4. The resulting plasmids 5 are designated pBTM- $\beta_2$ AR/AD4- $\beta$ ARK and pBTM-m2/AD4- $\beta$ ARK. respectively. PCR is used to amplify the DNA fragment containing bovine βarresting-1 (amino acids 1 to 437) [Lohse, et al., Science, 248:1547-1550 (1990)]. This fragment is inserted into pVP16 and is designated pVP16βarresting-1. PCR is used to amplify the DNA fragment containing rat βarresting-2 (amino acids 1 to 428) [Attramadal, et al., JBC, 267:17882-17890 (1992)] which is inserted into pVP16 to give plasmid pVP16-βarresting-2. A PCR strategy is also used to clone arresting into the pBTM116/AD4- $\beta$ ARK plasmid and the  $\beta$ AR and m2 fragments into pVP16. As above, the yeast split-hybrid YIDRM strain is transformed with the P-GR-arresting along 15 with peptide libraries (cloned into pRSURA3) or grown following transformation in the presence of combinatorial drug libraries.

Inhibitors identified in the split hybrid assay should effect disruption of protein/protein interaction either by: (i) inhibiting  $\beta$ ARK phosphorylation of the receptor, thus preventing recognition of the receptor by arresting, or (ii) by physical disruption of binding between the receptor and arresting. Agents that allow yeast growth for trivial reasons, *i.e.*, tetracycline analogues, can be easily identified through use of simple controls.

A first potential problem to overcome in this study is that cytoplasmic  $\beta$ ARK enzyme must be targeted to the substrate receptor and, once targeted, must phosphorylate the receptor at appropriate sites. In normal cells,  $\beta\gamma$  association serves to target  $\beta$ ARK to the cell membrane; the  $\beta$  subunit binds to both the  $\beta$ ARK PH domain and the isoprenylated  $\gamma$  subunit in association with the membrane. One possible means to encourage the necessary specific interactions is to target the binding components in the assay

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by tagging the proteins with nuclear localization signals, i.e., \( \beta ARK. \) the receptor cytoplasmic tail, and arresting, to the nucleus. The plasmids proposed for the study of the P-GR-arresting interaction all contain nuclear localization signal sequences adjacent to recombinant gene sequence.

A second problem is somewhat more difficult to approach. The current model is that receptors must be activated by ligand binding before being phosphorylated by  $\beta$ ARK, *i.e.*, targeting of  $\beta$ ARK via  $\beta\gamma$  is not sufficient for receptor phosphorylation. There are two possible explanations for this requirement. The first is that phosphorylation sites on the receptor are masked in the absence of ligand and ligand binding causes a conformational change which "unmasks" the phosphorylation sites. If this is the case, a fragment of the receptor containing the immediate phosphorylation site may be used as the  $\beta$ ARK target. However, although peptides representing portions of the  $\beta$ AR cytoplasmic tail can be phosphorylated by  $\beta$ ARK, the  $K_m$  for the phosphorylation reaction is poor, suggesting that the kinase may require some other part of the receptor for binding and that the unmasking of this binding site by agonist is a critical step.

This problem is addressed in two ways. In the first, the m2 muscarinic receptor is used in place of the  $\beta AR$  in view of previous results which indicate that the m2 protein is a good substrate for  $\beta ARK$ . The third cytoplasmic loop of the m2 receptor serves as both the binding site and phosphorylation site for kinase and which should allow use of a LexA/m2 receptor third cytoplasmic loop fusion gene as one component in the screening system.

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An alternative approach is to artificially mimic the activated state of the receptor. Haga, et al. U. Biol. Chem. 269:12594-12599 (1994)] have shown that the activity of βARK can be stimulated in vitro in the presence of mastoporan, a bee venom peptide. Mastoporan is believed to inimic the cytoplasmic face of an activated receptor and has been shown to increase the affinity of βARK for a GST-m2 receptor fusion protein by over

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four orders of magnitude. The same effect can be seen by using peptides representing the flanking regions of the m2 third cytoplasmic loop. Thus, mastoporan should also activate  $\beta$ ARK in the two-hybrid yeast strains, allow phosphorylation of the receptor fusion protein, and promote interaction with arresting. If mastoparan is needed, oligonucleotides containing the coding and non-coding nucleotide sequences of the 14-mer peptide (INLKALAALAKKIL-NH<sub>2</sub>, SEQ ID NO: 43) are annealed and ligated into prSADE2. The yeast split-hybrid strain YIDRM is transformed with pBTM- $\beta$ AR (or m2)/AD4- $\beta$ ARK, pVP16-arresting, pRSADE2-mastoparan, and a pRSURA3-peptide library or combinatorial drug library.

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Numerous modifications and variations in the invention as set forth in the above illustrative examples are expected to occur to those skilled in the art. Consequently only such limitations as appear in the appended claims should be placed on the invention.

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#### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
  - (i) APPLICANT: Hoekstra, Merl F.
  - (ii) TITLE OF INVENTION: Methods to Identify Compounds For Disrupting Protein/Protein Interactions
  - (iii) NUMBER OF SEQUENCES: 43
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      - (F) ZIP: 60606-6402
    - (v) COMPUTER READABLE FORM:
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      - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
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      - (C) TELEX: 25-3856
- (2) INFORMATION FOR SEQ ID NO:1:
  - - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs
      - (B) TYPE: nucleic acid

      - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
    - (ii) MOLECULE TYPE: DNA
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TTGGTGAGCG CTAGGAGTCA CTGCCAG

- (2) INFORMATION FOR SEQ ID NO:2:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 43 base pairs (B) TYPE: nucleic acid

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(ii) MOLECULE TYPE: DNA

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(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
TATACTCTAT CAATGATAGA GTAATTCATT ATGTGATAAT GCC	43
(2) INFORMATION FOR SEQ ID NO:3:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 42 base pairs  (B) TYPE: uncleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
ATTACTCTAT CATTGATAGA GTATATAAAG TAATGTGATT TC	42
(2) INFORMATION FOR SEQ ID NO:4:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
AATTCTGCTA GCCTCTGCAA AGC	23
(2) INFORMATION FOR SEQ ID NO:5:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRAMDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
CGCACGCGTC GAAGAAATCA CATTACTTTA TATA	34
(2) INFORMATION FOR SEQ ID NO:6:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
CGCACGCGTA TACTAAAAAA TGAGCAGGCA AG	3
(2) INFORMATION FOR SEQ ID NO:7:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 25 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
CGCGTACTCT ATCATTGATA GAGTA	25
(2) INFORMATION FOR SEQ ID NO:8:	-
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
ATGAGATAGT AACTATCTCA TGCGC	25
(2) INFORMATION FOR SEQ ID NO:9:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 48 base pairs (B) TYPS: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
CGCGTACTCT ATCATTGATA GAGTCTAGAC TCTATCAATG ATAGAGTA	48
(2) INFORMATION FOR SEQ ID NO:10:	•••
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
GCGACGCGTG CATGCCGTCT TCAAGAATTC CTCGAG	
CTCGAG	36

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(2) INFORMATION FOR SEQ ID NO:11:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
GCGACGCGTG CATGCCCACC GTACACGCCT ACTCGA	
(2) INFORMATION FOR SEQ ID NO:12:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 34 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
CATGGCATGC AAAAAAAAG AGTCATCCGC TAGG	3
(2) INFORMATION FOR SEQ ID NO:13:	_
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
CATGGCATGC TTAGCGATTG GCATTATCAC AT	3
(2) INFORMATION FOR SEQ ID NO:14:	3
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPS: nucleic acid (C) STRAMBEDNESS: single (D) TOPOLOSY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
TAATACGACT CACTATATAG GG	22
(2) INFORMATION FOR SEQ ID NO:15:	22
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

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(11) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
TCTAGACTTT GCCTTCGTTT ATC	2:
(2) INFORMATION FOR SEQ ID NO:16:	٤.
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 base pairs (B) TTPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
CGAAGGCAAA GATGTCTAGA TTAGATAAAA G	31
(2) INFORMATION FOR SEQ ID NO:17:	-
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 49 base pairs (B) TTPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
CGCGGATCCG CTTTCTCTTC TTTTTTGGAG ACCCACTTTC ACATTTAAG	49
(2) INFORMATION FOR SEQ ID NO:18:	.,
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH, 32 hase pairs  (B) TYPE: nu 32 hase pairs  (C) STRANDERS'S: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
AATTGCTCGA GTACTGTATG TACATACAGT AG	32
(2) INFORMATION FOR SEQ ID NO:19:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 32 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
ANTICTACTG TATGTACATA CAGTACTCGA GC	32

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(2) INF	ORMATION FOR SEQ ID NO:20:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDMESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:20:	
CCGGAATT	CT CGAGACATAT CCATATCTAA TC	32
(2) INFO	RMATION FOR SEQ ID NO:21:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 27 base pairs (B) TTPE: nucleic acid (C) STRANDEDINESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CCGGAATT	CA CTAATCGCAT TATCATC	27
(2) INFO	RMATION FOR SEQ ID NO:22:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 32 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:22:	
CATGCCAT	GG CCATGTCTAG ATTAGATAAA AG	32
(2) INFO	RMATION FOR SEQ ID NO:23:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:23:	
GCGAATTCC	C CAGGGCAACA GAATGCCACT	30
(2) INFOR	MATION FOR SEQ ID NO:24:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) (xi) (ccogaarii (2) INFO (ii) (xi) (cci) (xi) (xi) (xi) (xi) (xi) (xi) (xi) (x	(B) TTPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MoLECULE TYPE: DNA  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:  CCOGGAATTCT CGAGACATAT CCATATCTAA TC  (2) INFORMATION FOR SEQ ID NO:21:  (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:  CCCGGAATTCA CTAATCCCAT TATCATC  (2) INFORMATION FOR SEQ ID NO:22: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: DNA (xi) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: DNA (xi) SEQUENCE CEARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (C) TOPOLOGY: linear  (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23: (II) SEQUENCE DESCRIPTION: SEQ ID NO:23: (II) SEQUENCE DESCRIPTION: SEQ ID NO:23: (II) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (C) TOPOLOGY: linear  (II) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23: (II) SEQUENCE CHARACTERISTICS: (II) SEQUENCE CHARACTERI

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- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

# CGGGATCCTG GCTGGTTACC CAGGATGCCT TG

- (2) INFORMATION FOR SEQ ID NO:25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid

    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear

  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:
- CGCGGATCCG GATGACCATG GACTCTGGAG
- (2) INFORMATION FOR SEQ ID NO: 26:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 base pairs

    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26: CGCGGATCCT TAATCTGACT TGTGGCAGTA
- (2) INFORMATION FOR SEQ ID NO:27:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 32 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

CGCGGATCCC CATGACCATG GAATCTGGAG CC 32

- (2) INFORMATION FOR SEQ ID NO:28:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 31 base pairs
      - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
      - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28: CGCGGATCCG TGCTGCTTCT TCAGCAGGCT G
- (2) INFORMATION FOR SEQ ID NO:29:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 40 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

# ATGGTACCAG CGGCCGCTAG TCGTTTTACA ACGTCGTGAC

- (2) INFORMATION FOR SEQ ID NO:30:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 38 base pairs (B) TYPE: nucleic acid

      - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

ATGGTACCGC GGCCGCTTAT TTTTGACACC AGACCAAC

- (2) INFORMATION FOR SEQ ID NO:31:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 34 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

CGGAGATCTA AAGAGACTTT TCTCCGGAAC TCAG

- (2) INFORMATION FOR SEQ ID NO:32:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 29 base pairs
      - (B) TYPE: nucleic acid
      - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

CGGAGATCTT TACAGGAAGA CTGAACTGT

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- (2) INFORMATION FOR SEQ ID NO:33:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 29 base pairs
    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

CCACCGCGC AGTGCCAACC CCGATTTAC

- (2) INFORMATION FOR SEQ ID NO:34:
  - - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid
      - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA

  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

CATCCGCGGT GGTGATGGCA GGGGCTGA 28

- (2) INFORMATION FOR SEQ ID NO:35:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 28 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: Bingle (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

GGCTATCGAT ACGGCCCCCC CGACCGAT

- (2) INFORMATION FOR SEQ ID NO:36:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 28 base pairs
      - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
      - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

GCGTATCGAT CTACCCACCG TACTCGTC

- (2) INFORMATION FOR SEQ ID NO:37:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs

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- (B) TYPE: nucleic acid (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

CCTACTCTTA GGCCCGGGTC TTTTTAATGT ATCC

- (2) INFORMATION FOR SEQ ID NO:38:
  - (i) SEQUENCE CHARACTERISTICS:

    - (A) LENGTH: 18 base pairs
      (B) TYPE: nucleic acid
      (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

GGAATCACTA CAGGGATG

- (2) INFORMATION FOR SEQ ID NO:39:
  - (i) SEQUENCE CHARACTERISTICS:
    - ¡EQUENCE CHARACTERISTICS:
      (A) LENGTH: 1485 base pairs
      (B) TYPE: nucleic acid
      (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:	
ATGGACTTAA GAGTAGGAAG GAAATTTCGT ATTGGCAGGA AGATTGGGAG TGGTTCCTTT	6
GGTGACATTT ACCACGGCAC GAACTTAATT AGTGGTGAAG AAGTAGCCAT CAAGCTGGAA	120
TCGATCAGGT CCAGACATCC TCAATTGGAC TATGAGTCCC GCGTCTACAG ATACTTAAGC	180
GGTGGTGTGG GAATCCCGTT CATCAGATGG TTTGGCAGAG AGGGTGAATA TAATGCTATG	240
GTCATCGATC TTCTAGGCCC ATCTTTGGAA GATTTATTCA ACTACTGTCA CAGAAGGTTC	300
TCCTTTAAGA CGGTTATCAT GCTGGCTTTG CAAATGTTTT GCCGTATTCA GTATATACAT	360
GGAAGGTCGT TCATTCATAG AGATATCAAA CCAGACAACT TTTTAATGGG GGTAGGACGC	420
CGTGGTAGCA CCGTTCATGT TATTGATTTC GGTCTATCAA AGAAATACCG AGATTTCAAC	480
ACACATCGTC ATATTCCTTA CAGGGAGAAC AAGTCCTTGA CAGGTACAGC TCGTTATGCA	540
AGTGTCAATA CGCATCTTGG AATAGAGCAA AGTAGAAGAG ATGACTTAGA ATCACTAGGT	600
TATGTCTTGA TCTATTTTTG TAAGGGTTCT TTGCCATGGC AGGGTTTGAA AGCAACCACC	660
AAGAAACAAA AGTATGATCG TATCATGGAA AAGAAATTAA ACGTTAGCGT GGAAACTCTA	720
TGTTCAGGTT TACCATTAGA GTTTCAAGAA TATATGGCTT ACTGTAAGAA TTTGAAATTC	780
GATGAGAAGC CAGATTATTT GTTCTTGGCA AGGCTGTTTA AAGATCTGAG TATTAAACTA	840
GAGTATCACA ACGACCACTT GTTCGATTGG ACAATGTTGC GTTACACAAA GGCGATGGTG	900
GAGAAGCAAA GGGACCTCCT CATCGAAAAA GGTGATTTGA ACGCAAATAG CAATGCAGCA	960
AGTGCAAGTA ACAGCACAGA CAACAAGTCT GAAACTTTCA ACAAGATTAA ACTGTTAGCC	1020
ATGAAGAAAT TCCCCACCCA TTTCCACTAT TACAAGAATG AAGACAAACA TAATCCTTCA	1080
CCAGAAGAGA TCAAACAACA AACTATCTTG AATAATAATG CAGCCTCTTC TTTACCAGAG	1140
GAATTATTGA ACGCACTAGA TAAAGGTATG GAAAACTTGA GACAACAGCA GCCGCAGCAG	1200
CAGGTCCAAA GTTCGCAGCC ACAACCACAG CCCCAACAGC TACAGCAGCA ACCAAATGGC	1260
CAAAGACCAA ATTATTATCC TGAACCGTTA CTACAGCAGC AACAAAGAGA TTCTCAGGAG	1320
CARCAGCAGC ARGTTCCGAT GGCTACAACC AGGGCTACTC AGTATCCCCC ACAAATAAAC	1380
AGCARTARTT TTARTACTAR TCARGCATCT GTACCTCCAC ARATGAGATC TARTCCACAA CAGCCGCCTC ARGATARACC AGCTGGCCAG TCARTTTGGT TGTAR	1440
(2) INFORMATION FOR SEQ ID NO:40:	1485
TOR DEQ ID NO:40:	

<sup>(</sup>i) SEQUENCE CHARACTERISTICS:

SEQUENCE CHARACTERISTICS:

(A) LENGTH: 2625 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

<sup>(</sup>ii) MOLECULE TYPE: DNA

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(ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 796..2580

(xi) SEQUENCE DESCRIPTION: SEO ID NO:40: CATTITCITA ATTCTTTAT GIGCTITTAC TACTITGTIT AGTICAAAAC AATAGTCGTT 60 ATTCTTAGGT ACTATAGCAT AAGACAAGAA AAGAAAAATA AGGGACAAAT AACATTAGCA 120 GAAGTACGGT ATATTTTACT GTTACTTATA TACTTTCAAG AAGATGAGTT AAATCGGTAG 180 CCAGTGTAGA AAAATAATAA TAAGGGTCAT CGATCCTTCG CATTTTATTA TCCAATTAAA 240 GATACGAATC ACGGCAAACT ATATTCAAAG CTCATAGATA ATCGTCGTAA GGCTGACACT 300 GCAGAAGAAA AGTCATAATT TGAATACTAG CCGGTATGAA ACTGTGATTG ATTAACCTGG 360 GGTTACCTAA AGAGAACATA AGTAATACTC ATGACAGAAT CAAAACACAA TACAAAATTT 420 ATCCGAACCT CGGCCCGACT GCGGCTCGCC GGGAAAGGGG ACAACCGCTT CTATCCGTCG 480 ACTAACTTCA TCGGCCCAAT GGAAGCTATG ATATGGGGAT TTCCATTGAG CCGATAGCAA 540 TGTAGGGTAA TACTGTTGCG TATATAGTGA TAGTTATTGA ATTTTATTAC CCTGCGGGAA 600 TATTGAGACA TCACTAAGCA CGAATTTTAC GTCTGAGGAA AGTTGAATGA TGGCCAAATA 660 ACCAGGAAAA ACAAATATTG AATCCTTGTG AAGGATTCCA CAGTTGTTTA ATCCTCCTTA 720 AGCTCACTTA GTATCAATTG TCTAAATAAT ATTGCTTTGA ATCTGAAAAA AATAAAAGTA 780 CCTTCGCATT AGACA ATG TCA CTG CCG CTA CGA CAC GCA TTG GAG AAC GTT Met Ser Leu Pro Leu Arg His Ala Leu Glu Asn Val ACT TCT GTT GAT AGA ATT TTA GAG GAC TTA TTA GTA CGT TTT ATT ATA 879 Thr Ser Val Asp Arg Ile Leu Glu Asp Leu Leu Val Arg Phe Ile Ile AAT TGT CCG AAT GAA GAT TTA TCG AGT GTC GAG AGA GAG TTA TTT CAT Asn Cys Pro Asn Glu Asp Leu Ser Ser Val Glu Arg Glu Leu Phe His 927 TTT GAA GAA GCC TCA TGG TTT TAC ACG GAT TTC ATC AAA TTG ATG AAT Phe Glu Glu Ala Ser Trp Phe Tyr Thr Asp Phe Ile Lys Leu Met Asn 975 CCA ACT TTA CCC TCC CTA AAG ATT AAA TCA TTT GCT CAA TTG ATC ATA Pro Thr Leu Pro Ser Leu Lys Ile Lys Ser Phe Ala Gln Leu Ile Ile 1023 AAA CTA TGT CCT CTG GTT TGG AAA TGG GAC ATA AGA GTG GAT GAG GCA 1071 Lys Leu Cys Pro Leu Val Trp Lys Trp Asp Ile Arg Val Asp Glu Ala CTC CAG CAA TTC TCC AAG TAT AAG AAA AGT ATA CCG GTG AGG GGC GCT Leu Gln Gln Phe Ser Lys Tyr Lys Lys Ser Ile Pro Val Arg Gly Ala 1119 GCC ATA TIT AAC GAG AAC CTG AGT AAA ATT TTA TTG GTA CAG GGT ACT 1167

Ala Ile Phe Asn Glu Asn Leu Ser Lys Ile Leu Leu Val Gln Gly Thr

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12	5	. Aai	J 361	Let	130	Pne	Pro	Arc	Gly	135	Ile	e Ser	Lys	s Asp	GAA Glu 140	1215
			,	145	Cya	, ile	Arg	GIU	150	Lys	Glu	: Glu	Ile	: Gly		126
			160	-,-		- Aug	veb	165	GIN	Pne	11e	Glu	170	Asn	ATT	1311
	,	175		.,.	-yo	116	180	Leu	ire	Ser	Gly	Val 185	Ser	Glu	GTC Val	1359
	190		-,-		01	195	Arg	Asn	GIU	ire	200	Lys	Ile	Glu	TGG	1407
205	•			-,-	210	UCL	Lys	1111	mec	215	rys	Ser	Asn	Ile	Lys 220	1455
-	•			225	-	Mec	nec	Arg	230	Leu	Ser	Met	Trp	Leu 235		1503
			240	***	Lys	ABII	GIU	245	Gin	Leu	Lys	Ser	Tyr 250	Ala	GAA Glu	1551
		255					260	116	mr	rys	GIU	265	Gin	Ile	Asp	1599
	270		GAG Glu			275	nec	rea	HIS	rnr	280	Val	Gln	Ala	Asn	1647
285			AAT Asn		290	Ser	ABIL	Gly	GIn	295	Pro	Ser	Ser	Gln	Glu 300	1695
			TTG Leu	305			561	Gry	310	nıs	Asn	Gin	Gln	Lys 315	Asp	1743
			TCG Ser 320				G.1.1	325	GIN	PIO	ser	He	Phe 330	Pro	Ser	1791
CTT Leu	TCT Ser	GAA Glu 335	CCG Pro	TTT Phe	GCT Ala		AAT Asn 340	AAG Lys	AAT Asn	GTT Val	ATA Ile	CCA Pro 345	CCT Pro	ACT Thr	ATG Met	1839
	350		AAC Asn			355	Jer	Abii	Pro	Gin	360	Phe	Ala	Thr	Met	1887
AAT	GGC	CAG	CCT	TTT	GCA	CCT	TTC	CCA	TTT	ATG	TTA	CCA	TTA	ACT	AAC	1935

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		41	5				420	FIL	Pne	ser	Leu	425	Pro	Ala	CCT Pro	2079
	430	)				435			361	Jei	440	GIY	Gin	Leu	TTA Leu	2127
445					450			Д	361	455	vai	GIN	Ser	Ser	460	2175
				465		TTA Leu	٠	ALY	470	inr	Asp	Leu	Asn	Ser 475	Leu	2223
			480	)		GAA Glu		485	nis	ser	Asn	Ser	G1n 490	Ala	Leu	2271
		495		•	-,-	CCA Pro	500	Ser	ser	Gin	Lys	505	His	Ala	Ser	2319
	510					TTA Leu 515		74011	veh	ser	520	ser	Gly	He	Gln	2367
525					530	TTC Phe			061	535	мвр	GIU	GIu	Val	G1u	2415
				545	-	AAT Asn			550	vai .	мър	iie .	GIY	Va1 555	Asn	2463
			560		-	GAC .		565	ALY .	Ser (	JIN ,	Lys	51u 570	Lys	Pro	2511
		575			•		580	ocu ,		nia 3	er /	ъца ( 585	ilu :	Ser	Asn	2559
AGT (	TA ( /al (	GAA Glu	TGG Trp	GGG Gly		GGG 1 Gly 595	TAAA?	стто	CA CO	стсс	GACT	r TCJ	AGAG'	ГААС		2610
ACAG	ATC	CA C	AGTA													2625

	(2)	INFORMATION	FOR	SEO	ΤD	NO - 41
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- (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 6854 base pairs
  - (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA

### (ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 2050..4053

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

AGCTTCTCCC TTTTCCTTCA GTGCTGCTAC TCTCTGCTCT CCACTTAAGT GTTACAATTA 60 ATTTGCAGCT AGTTTGCAGT TCGTACAACC TCGCCTATTC TTGTAACGAA GAAGAACGTA 120 TTTATAATAT TGGGCTGTAA TGTGTTGAGT TTAGTAATAG ATAAAGTAGG ACAGAGTTCT 180 GTCTTTGTTT ATCTATGGGG TTCAGAGTGA TAAGGGGCAG GATAAGGAAG TTAAAAAAAA 240 AAAGGTTACG TTATATAACG AAAGAAAAGA AACGAGCGAA GTGCCAACTA TAGCCCAATA 300 TCAAGAATGC AAGTCAGCAA AGTACAGTAA TCGTATGAAG ATACGCGATG CGTAATATCC 360 CTCAAGGGCT CCGGATCAGA AAAGCTAAGG GAAGATCCTT ACATTACACG GCGTGCGACA 420 GACTCGAACC ACAGCTAACT TCTCGTGAAA AGATGGCTTC AACTTCGCTC TTGCAATAAC 480 TTTGAAACAC ACGAACAAAG GTTTATTGCG CTTGATTAAC GTTGGAAGTA TATGATACTA 540 ATACTACTTT GTTCTCTAAG TCATCGCTAT ATGTTTATCT CGAGGAAAAG GTGCACGGCG 600 GTACACAATT ACTTCGCCGT TTCGGGTAAA ACAAGTGTTA CATTTATAAT ATATATGTAT 660 ATATGTATGT GCGCGTAAGT ATATGCCGTT CATAACAAAT CATCTTCTTG TTGCTGGATG 720 GACTCCTTAA TTTTATTCAA AATGGTAATT TTCCATTTAT CTAGTCTCAT AAAATTGTCA 780 AACTECTTAC AGTGTTCGCT TAGCTGCTCG CTATCACCTT CATTAACAGC ATCGATTAAA 840 CTTTTCAAGA AATTTGACTC CCTTGAATCC GCAAAATTCG GATCTTCACT TTGACCCTCT 900 TGTAAAGTTC TTGCAGCAGC GACTGCATCA GTAGCAGCTA GCTGACAAAG CCCTTTTTTT 960 AGGAAGTAAT CCTTCAAACT CCATTGGCTC AATCTATTGC CCATGCTGCT CTTGATCAAC 1020 TTCGAATATA TATCACTTGC TTCAATATAT TGACCGTCAA GAGCCTTTAG ATCTGCGCAT 1080 TTGATAAAAC ACTTATTCGA TAATGCTACC GACTGGTCTT GGGCATACCA CTCACCAGCG 1140 AGCTCATAGC AATCTATAGC TTTTGCATAG TCATGCAAAT CATTTTCTAG AATTTCTCCA 1200 AGCTCAAACT TGAAATTAGC ACCTCTCCGG AACTGCCCCC TATGAGTAAA AATTTGAATA 1260 GCATTTTCTA ATGAATCCAC GGCGTTCACA GAGTTTCCAC CGCTTTTAAA GCATTTATAA 1320 GCCTCTACGT AGGTATTTCC TGCTTCGTCT TCATTACCAG CCTTTTTCTG ATAGTCAGCA 1380

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COMMONANA	
GCTTTCAAAA ACGAGTCTCC TGCCAAGTTT AACTCTTTTC TTAGACGGTA AATGGTGG	
GCTTGGACAC AAAGATCAGC AGCCTCCTCA AACTTGTATG AATCAGAACC GCTAAACAA	
TTCATGAAAC CCGATGAAGG AACACCCTTC TTCTCAGCCT TAACACAACG GGAAATATC	CA 1560
ATTCCCGTAT TTCAATGTTA GTAATTTGCC TTCGTAAATT ACGGAATCAC ATAGCTTTC	CA 1620
TTTTGTTCCT TTGATATATT TCCCTACTAC ATACTCTTTT CAATAACTCT ACAGGGTCT	rG 1680
ACATTITTAA CTTTCAGGTT AATGATGGTG TTCTTACTAT ATTCTCGAGT CGTACAGAA	AG 1740
TTAGTTCAGA TAAACTGCTT CGGTGCTGCC CACTTCTTAT CATTACTTCA ACTITACCT	T 1800
CCCTATACCT GTGTGTCCTT ATTAATTCAA GTTAATCCGA GGTAATAGAT TAGGGTAAC	C 1860
TTCAATGATG TCACGAAACA CGGATGCTGC AACTTTGCGA TTTTTTCCTG GAAAAGAAT	`A 1920
ACAATTAAAG GCAGCCTTTC AGCTGAGATT ACCAGCAGGT CTTTGGAGAT TAGCGCAAG	A 1980
AGAAGTGTGA TATAGTACTC ATAGAGGCAG GCTACAGACT AGGGAAAGCG TGTTCAACA	A 2040
CAATAAGAA ATG GAG ACC AGT TCT TTT CAG AAT COT COT COT	2088
1 5 50 Ash Ala Pro Pro Ala Ala	
ATC AAT GAT GCT CAG GAT AAT AAT ATA AAT ACG GAG ACT AAT GAC CAG	2136
Ile Asn Asp Ala Gln Asp Asn Asn Ile Asn Thr Glu Thr Asn Asp Gln  15  20  25	-150
GAA ACA AAT CAG CAA TCT ATC GAA ACT AGA GAT GCA ATT GAC AAA GAA	2184
Glu Thr Asn Gln Gln Ser Ile Glu Thr Aga GAT GCA ATT GAC AAA GAA 30 35 40 45	
AAC GGT GTG CAA ACG GAA ACT GGT GAG AAC TCT GCA AAA AAT GCC GAA	2232
Asn Gly Val Gln Thr Glu Thr Gly Glu Asn Ser Ala Lys Asn Ala Glu  50  55  60	2232
CAA AAC GIT TOT TOT ACA AND THE	2280
Sin Asn Val Ser Ser Thr Asn Leu Asn Asn Ala Pro Thr Asn GGT GCT  65  70  75	2280
FTG GAC GAT GAT GTT ATC GCA AND	
Leu Asp Asp Asp Val Ile Pro Asn Ala Ile Val Ile Lys Asn Ile Pro	2328
TTT GCT ATT AND AND CAG GAR	
95 Sed Asp Ite Ite Glu Glu Met Asp	2376
TT CCC CTT CCT TAT CCC TTO	
10 Ash Tyr his Phe Asp Ash Gly Ile Phe	2424
GA GGA CTA GCC TTT GCC AND THE COL	
GA GGA CTA GCC TTT GCG AAT TTC ACC ACT CCT GAA GAA ACT ACT CAA rg Gly Leu Ala Phe Ala Asn Phe Thr Thr Pro Glu Glu Thr Thr Gln	2472
135 140	
TG ATA ACT TCT TTG AAT GGA AAG GAA ATC AGC GGG AGG AAA TTG AAA al lie Thr Ser Leu Asn Gly Lys Glu Ile Ser Gly Arg Lys Leu Lys	2520
145 150 150 155 Let Lyg	

GTG Val	GAA Glu	TAT Tyr 160	AAA Lys	AAA Lys	ATG Met	CTT Leu	CCC Pro 165	CAA Gln	GCT Ala	GAA Glu	AGA Arg	GAA Glu 170	AGA Arg	ATC Ile	GAG Glu	2568
AGG Arg	GAG Glu 175	AAG Lys	AGA Arg	GAG Glu	AAA Lys	AGA Arg 180	GGA Gly	CAA Gln	TTA Leu	GAA Glu	GAA Glu 185	CAA Gln	CAC His	AGA Arg	TCG Ser	2616
TCA Ser 190	TCT Ser	AAT Asn	CTT	TCT Ser	TTG Leu 195	GAT Asp	TCT Ser	TTA Leu	TCT Ser	AAA Lys 200	ATG Met	AGT Ser	GGA Gly	AGC Ser	GGA Gly 205	2664
AAC Asn	AAT Asn	AAT Asn	ACT Thr	TCT Ser 210	AAC Asn	AAT Asn	CAA Gln	TTA Leu	TTC Phe 215	TCG Ser	ACT Thr	CTA Leu	ATG Met	AAC Asn 220	GGC Gly	2712
ATT Ile	AAT Asn	GCT Ala	AAT Asn 225	AGC Ser	ATG Met	ATG Met	AAC Aan	AGT Ser 230	CCA Pro	ATG Met	AAT Asn	AAT Asn	ACC Thr 235	ATT Ile	AAC Asn	2760
AAT Asn	AAC Asn	AGT Ser 240	TCT Ser	AAT Asn	AAC Asn	AAC Asn	AAT Asn 245	AGT Ser	GGT Gly	AAC Asn	ATC Ile	ATT Ile 250	CTG Leu	AAC Asn	CAA Gln	2808
Pro	TCA Ser 255	CTT Leu	TCT Ser	GCC Ala	CAA Gln	CAT His 260	ACT Thr	TCT Ser	TCA Ser	TCG Ser	TTG Leu 265	TAC Tyr	CAA Gln	ACA Thr	AAC Asn	2856
GTT Val 270	AAT Asn	AAT Asn	CAA Gln	GCC Ala	CAG Gln 275	ATG Met	TCC Ser	ACT Thr	GAG Glu	AGA Arg 280	TTT Phe	TAT Tyr	GCG Ala	CCT Pro	TTA Leu 285	2904
CCA Pro	TCA Ser	ACT Thr	TCC Ser	ACT Thr 290	TTG Leu	CCT Pro	CTC Leu	CCA Pro	CCC Pro 295	CAA Gln	CAA Gln	CTG Leu	GAC Asp	TTC Phe 300	AAT Asn	2952
GAC Asp	CCT Pro	GAC Asp	ACT Thr 305	TTG Leu	GAA Glu	ATT Ile	TAT Tyr	TCC Ser 310	CAA Gln	TTA Leu	TTG Leu	TTA Leu	TTT Phe 315	AAG Lys	GAT Asp	3000
AGA Arg	GAA Glu	AAG Lys 320	TAT Tyr	TAT Tyr	TAC Tyr	GAG Glu	TTG Leu 325	GCT Ala	TAT Tyr	CCC Pro	ATG Met	GGT Gly 330	ATA Ile	TCC Ser	GCT Ala	3048
TCC Ser	CAC His 335	AAG Lys	AGA Arg	ATT Ile	ATC Ile	AAT Asn 340	GTT Val	TTG Leu	TGC Cys	TCG Ser	TAC Tyr 345	TTA Leu	GGG Gly	CTA Leu	GTA Val	3096
GAA Glu 350	GTA Val	TAT Tyr	GAT QaA	CCA Pro	AGA Arg 355	TTT Phe	ATT Ile	ATT Ile	ATC Ile	AGA Arg 360	AGA Arg	AAG Lys	ATT Ile	CTG Leu	GAT Asp 365	3144
CAT His	GCT Ala	AAT Asn	TTA Leu	CAA Gln 370	TCT Ser	CAT His	TTG Leu	CAA Gln	CAA Gln 375	CAA Gln	GGT Gly	CAA Gln	ATG Met	ACA Thr 380	TCT Ser	3192
GCT Ala	CAT His	CCT Pro	TTG Leu 385	CAG Gln	CCA Pro	AAC Asn	TCC Ser	ACT Thr 390	GGC Gly	GGC Gly	TCC Ser	ATG Met	AAT Asn 395	AGG Arg	TCA Ser	3240
CAA Gln	TCT Ser	TAT Tyr 400	ACA Thr	AGT Ser	TTG Leu	TTA Leu	CAG Gln 405	GCC Ala	CAT His	GCA Ala	GCA Ala	GCT Ala 410	GCA Ala	GCG Ala	AAT Asn	3288

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AGT Se	Ile 41	e Se:	C AAT	CAC Glr	G GCG	Val	. Asr	AA :	TCT Ser	TC(	AAG ASI 425	s Se i	AAT Ası	Thi	Ile	3336
AA( A61 43(	se:	AA 1	AA C	GGI	AAC ASI 435	ı Gly	AAC	AAT Asr	GTC Val	Ile 440	: Ile	CAA 7	AAC Asr	AAT Asr	AGC Ser 445	3384
GC(	Se	C TC	A ACA	Pro 450	· Lyε	ATT Ile	Ser	Ser	Gln 455	Gly	CA#	TTC Phe	TCC Ser	Met 460	CAA Gln	3432
Pro	ACJ Thi	Let	ACC Thr 465	: Ser	Pro	Lys	Met	AAC Asn 470	Ile	CAC	CAT	AGT Ser	Ser 475	Gln	TAC	3480
AAT	Ser	GC/ Ala 480	і Абр	Gln	Pro	CAA Gln	CAA Gln 485	Pro	CAA Gln	Pro	CAA Gln	ACA Thr 490	Gln	CAA Gln	AAT Asn	3528
Val	Glr 495	ı ser	GCT	GCG	Gln	Gln 500	Gln	CAA Gln	TCT Ser	TTT	Leu 505	Arg	CAA Gln	CAA Gln	GCT Ala	3576
ACT Thr 510	Leu	ACA Thr	Pro	Ser	Ser 515	Arg	ATT	CCA Pro	TCC	GGT Gly 520	Tyr	Ser	GCC	AAC	CAT His 525	3624
TAT	Gln	ATC	AAT Asn	Ser 530	val	AAT Asn	CCC	TTA Leu	CTG Leu 535	AGA Arg	AAT Asn	TCT Ser	CAA Gln	ATT Ile 540	TCA Ser	3672
Pro	Pro	AAT Asn	Ser 545	CAA Gln	ATC	CCA Pro	ATC Ile	AAC Asn 550	Ser	CAA Gln	ACC Thr	CTA Leu	TCC Ser 555	CAA Gln	GCG Ala	3720
GIn	Pro	560	Ala	Gln	Ser	Gln	Thr 565	Gln	Gln	Arg	Val	CCA Pro 570	Val	Ala	Tyr	3768
GIII	575	MIG	Ser	Leu	ser	580	Gin	Gln	Leu	Tyr	<b>A</b> sn <b>5</b> 85	CTT Leu	Asn	Gly	Pro	3816
590	ser	АТА	Asn	Ser	595	Ser	Gln	Leu	Leu	Pro 600	Gln	CAC His	Thr	Asn	Gly 605	3864
TCA Ser	GTA Val	CAT His	TCT Ser	AAT Asn 610	TTC Phe	TCA Ser	TAT Tyr	CAG Gln	TCT Ser 615	TAT Tyr	CAC His	GAT Asp	GAG Glu	TCC Ser 620	ATG Met	3912
TTG Leu	TCC Ser	GCA Ala	CAC His 625	AAT Asn	TTG Leu	TAA naA	AGT Ser	GCC Ala 630	GAC Asp	TTG Leu	ATC Ile	TAT Tyr	AAA Lys 635	TCT Ser	TTG Leu	3960
AGT Ser	CAC His	TCT Ser 640	GGA Gly	CTA Leu	GAT Asp	Asp	GGC Gly 645	TTG Leu	GAA Glu	CAG Gln	GGC Gly	TTG Leu 650	AAT Asn	CGT Arg	TCT Ser	4008
Leu	AGC Ser 655	GGA Gly	CTG Leu	GAT Asp	TTA Leu	CAA Gln 660	AAC Asn	CAA Gln	AAC Asn	AAG Lys	AAG Lys 665	AAT Asn	CTA Leu	TGG Trp		4053

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TAATATATAC	TICCATTAIT	CIAIGAIIAI	AGAGIIIGII	IGGIAITIGI	ATATCOCACO	4113
ATACAAGTAA	TGAGGGGTGC	TTACACAAGA	TAAAAGATAA	<b>ТАТАТАБАА</b>	ATATATATA	4173
AAAACCATCA	AAAACACCAT	TGAAAAAAA	TATAAAAAAA	АЛАААААТА	ACCGAATATG	4233
aatatgaaat	TAATGATCAT	GATGAAGTTA	ATTTTTACTG	AGAAACGTCA	CCTAATGTCG	4293
ATGAAACGAT	GATAATGAAT	GAATGATGAG	GCTACTITAA	GTAACGCAAT	GTAATCAAGC	4353
CAAAATTATC	CCTCTTTTT	TTTTTTCCCT	CTTTTGAGAT	ATTTTTTTT	ACCTACTACT	4413
TACTTTTTT	TTTTGAACGT	TCTTTTCCCA	CATACTTTTA	TATATGGTAT	TTATATGTAC	4473
GATGTTTAAT	CACAGAGATG	TTTCTACCTT	ACTCGATATT	GTTTTTGCAT	TAATTGATAT	4533
CTTGCTCACT	GCATCATTGG	CGGTATTTGT	AGTATATAGA	AAGTCGGGTA	ACAATAATT::	4593
ATTGACATTT	CTTTGTTTAC	AATGATCAGA	GAAGAGCAGA	AAGTTTCATA	GTCAAACGT"."	4653
CAGGCCAATT	GAACAAGAAA	TTATTCGTTT	TTTTAGTCGT	TGAGTGTTCA	ACTGACATGC	4713
TATTTTGGTG	GITCTTGATT	AATTGGGGGC	TTCATTGTTT	GAAATAAAGA	GTCGGGAAAA	4773
TAGCACAGAA	ACAAAGCATA	TTAAAAGAGG	CAAAAGAAGA	AAGAACGAAT	ataaaaggta	4833
AAAAAGGAAA	AGCATTGCTA	TTCTTTTCTC	ATAGGTGTTA	TTCATACCGC	CCTCTCTCTT	4893
CTTCCTTCTT	CATTAATTAG	TCTCCGTATA	ATTTGCAGAT	AATGTCATTA	ACAGCAAACG	4953
ACGAATCGCC	AAAACCCAAA	AAAAATGCAT	TATTGAAAAA	CTTAGAGATC	GATGATCTGA	5013
TACATTCTCA	ATTTGTCAGA	AGCGATACAA	ATGGACATAG	AACTACAAGA	CGACTATTCA	5073
ACTCCGATGC	CAGTATATCA	CATCGAATAA	GAGGAAGTGT	TCGGTCTGAT	AAAGGCCTTA	5133
AAAAAATA	AAAAGGGTTG	ATTTCCCAGC	AGTCCAAACT	TGCGTCAGAA	AATTCTTCTC:	519
AAAATATCGT	TAATAGGGAC	AATAAGATGG	GAGCAGTAAG	TTTCCCCATT	ATTGAACCTA	525
ATATTGAAGT	CAGCGAGGAG	TTGAAGGTTA	GAATTAAGTA	TGATTCTATC	AAATTTTTCL	531
ATTTTGAAAC	ACTANTATCT	AAATCTTCAG	TCATAGCACC	TTTAGTTAAC	AATATAAAA	537
CATCATCCGC	TCCTCTAATC	GGGTTTCAAA	GAAGAGTTAA	CAGGTTAAAG	CAAACATGGG	543
ATCTAGCAAG	CGAAAACATG	GAGTACCCAT	ATTCTTCTGA	TAATACGCCA	TTCAGGGATA.	549
ACGATTCTTC	GCAATGGTAC	GTACCATACG	GCGGAACAAT	TAAAAAAAA '	AAAGATTTCA.	555
GTACAAAAA	AACTTTACCC	ACCTGGGAAG	ATAAAATA S	GTTTCTTAC	TTTTTAGAAA.	561
ACTCTAAGT	TGCAACGTAC	ATTAATGGTA	ACGTATCACT	TTGCAATCAT	AATGAAACCG	567
ATCAAGAAA	CGAAGATAGG	AAAAAAAGG/	AAGGGAAAGT	ACCAAGAATO	AAAAATAAAG	573
TGTGGTTTT	CCAGATAGAA	TACATTGTT	TTCGAAATT	TGAAATTAA	CCTTGGTATA.	579
CATCTCCTT	r TCCGGAACAC	ATCAACCAA	ATAAAATGGT	TTTTATATG	GAGTTCTGCC	585
TAAAATATA	GACTTCTCG	TATACTTTT	T ATAGACACCA	ACTAAAGTG	CTAACTTTTA	591
AGCCCCCCG	AAATGAAATT	TATCGCGACG	GTAAGCTGT	TGTTTGGGA	ATTGATGGGC	597

6033

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AGACTTTGTA TTACGATGTT GAACCGTTTA TATTCTATAT TCTAACGGAG AGAGAGGATA 6093

GGGAGAATGT CTTGTATTGT CAAAATCTTT GCCTGTTGGC AAAATGTTTT ATCAATTCTA

CAGAGAACCA TCCCTATCAA AACGCAGCCA AATTCCATTT CGTAGGCTAT TTCTCCAAGG	6153
AAAAATTCAA CTCCAATGAC TATAACCTAA GTTGTATTTT AACTCTACCC ATATACCAGA	6213
GGAAAGGATA TGGTCAGTTT TTGATGGAAT TTTCATATTT ATTATCCAGA AAGGAGTCAA	6273
AATTTGGAAC TCCTGAAAAA CCATTGTCGG ATTTAGGATT ATTGACTTAC AGAACGTTTT	6333
GGAAGATAAA ATGTGCTGAA GTGCTATTAA AATTAAGAGA CAGTGCTAGA CGTCGATCAA	6393
ATAATAAAA TGAAGATACT TITCAGCAGG TTAGCCTAAA CGATATCGCT AAACTAACAG	6453
GAATGATACC AACAGACGTT GTGTTTGGAT TGGAACAACT TCAAGTTTTG TATCGCCATA	6513
AAACACGCTC ATTATCCAGT TIGGATGATT TCAACTATAT TATTAAAATC GATTCTTGGA	6573
ACAGGATTGA AAATATTTAC AAAACTTGGA GCTCAAAAAA CTATCCTCGC GTCAAATATG	6633
ACAAACTATT GTGGGAACCT ATTATATTAG GGCCGTCATT TGGTATAAAT GGGATGATGA	6693
ACTTAGAACC CACCGCATTA GCGGACGAAG CTCTTACAAA TGAAACTATG GCTCCGGTAA	6753
TTTCGARTAA CACACATATA GAAAACTATA ACAACAGTAG AGCACATAAT AAACGCAGAA	6813
GAAGAAGAAG AAGAAGTAGT GAGCACAAAA CATCCAAGCT T	6854
(2) INFORMATION FOR SEQ ID NO:42:	
(A) LENGTH: 2814 base pairs (B) TYPE: nucleic acid (C) STRANDENNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1.696	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
AA THC CAA TAC ACC AAA CAG CTG CAT THC CCT GTG GGG CCC AAA TCC llu Phe Gln Tyr Thr Lys Gln Leu His Phe Pro Val Gly Pro Lys Ser 1 10 15	48
CA AAC TOT GAG GTA GCG GAA ATT CTT TTA CAC TGC GAC TGG GAA AGG hr Asn Cys Glu Val Ala Glu Ile Leu His Cys Asp Trp Glu Arg $\begin{array}{cccccccccccccccccccccccccccccccccccc$	96
AC ATA AAT GTT TTA AGT ATA ACA AGA ACA CCA AAT GTT CCT AGT GGT yr 18 Asn Val Leu Ser Ile Thr Arg Thr Pro Asn Val Pro Ser Gly 45 45	144
CC AGT TTC AGC ACC AGA ACG AGG TAC ATG TTC CGA TGG GAT GAC CAG hr Ser Phe Ser Thr Arg Thr Arg Tyr Met Phe Arg Trp Asp Asp Gln	192

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6	5	Gly	٠,٠		70	Dyb	116	Ser	Pne	75 75	Val	Asp	Trp	Asn	Ala 80		240
Se	C AGT	Trp	Ile	Lys 85		ATG Met	GTA Val	GAG Glu	AGC Ser 90	Asn	TGT	AAA Lys	AAT Asn	GGA Gly 95	Gln		288
		GCC	100	<i>_</i>	ASP	Deu	vai	105	Leu	Val	Glu	Glu	Phe 110	Val	Glu		336
AA. Ly	A TAC	Val 115	GAA Glu	TTG	AGC Ser	AAA Lys	GAA Glu 120	AAA Lys	GCA Ala	GAT Asp	ACA Thr	CTC Leu 125	AAG Lys	CCG Pro	TTG Leu		384
Pre	Ser 130	GTT Val	ACA Thr	TCT	TTT Phe	GGA Gly 135	TCA Ser	CCT Pro	AGG Arg	AAA Lys	GTG Val 140	GCA Ala	GCA Ala	CCG Pro	GAG Glu		432
Let 145	S TCG Ser	Met	GTA Val	CAG Gln	CCG Pro 150	GAG Glu	TCG Ser	AAA Lys	CCA Pro	GAA Glu 155	GCT Ala	GAG Glu	GCG Ala	GAA Glu	ATC Ile 160		480
TC) Sei	GAA Glu	ATA Ile	GGC Gly	AGC Ser 165	GAC Asp	AGA Arg	TGG Trp	AGG Arg	TTT Phe 170	AAC Asn	TGG Trp	GTG Val	AAC Asn	ATA Ile 175	ATA 11e		528
ATC Ile	: TTG : Leu	GTG Val	CTC Leu 180	TTG Leu	GTG Val	TTA Leu	AAT Asn	CTG Leu 185	CTG Leu	TAT Tyr	TTA Leu	ATG Met	AAG Lys 190	TTG Leu	AAC Asn		576
Lys	Lys	ATG Met 195	GAT Asp	AAG Lys	CTG Leu	****	AAC Asn 200	CTC Leu	ATG Met	ACC Thr	CAC His	AAG Lys 205	GAC Asp	GAA Glu	GTT Val		624
GTA Val	GCG Ala 210	CAC His	GCG Ala	ACT Thr	CTA Leu	TTG Leu 215	GAC Asp	ATA Ile	CCA Pro	Ala	CAA Gln 220	GTA Val	CAA Gln	TGG Trp	TCA Ser		672
AGA Arg 225		AGA Arg	AGG Arg	GGA Gly	GAC Asp 230	GTG Val	TTG Leu	TAAC	AGAG	TA A	TCAT	GTAA	TA T	TGTA	TGTA		726
AGG	TTATO	та т	GTTC	GTAT	G GT	ATGG	AAAA	AAA	алал	AAA	AAAG	GATG	CT A	TGTG	GAGAA		786
															GCCTG		846
															GAAAT		906
															ACCTG		966
															GTTTG		1026
															ACCGC		1086
															GACGA	:	1146
															AATAA	:	1206
															GCACA	:	1266
CIG	GCCT	TT TO	CAA	TATG	C CG	ATGC	TOT	TTT	'AGG	CAA A	ATTC	CAA:	rg g	CATC	CCGG	- 1	1326

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ATTGAAGCGA TAATCCTTAG CGTCCTGAAC CAATTGCTTA CTAGACTTCA TGACCTACCG	1386
GGGCCAGATA AAGATGCGGA AGGAAGAGAA AAAATGTATA GTGGTTGGTG AACCGCAACA	1446
ATAATTCGTG CCAACACTTT AATCGAAGCA AAAATTGTCT TGTATGTTAT TAATATTATC	1506
TATCTAACCA TTGATTTACG TATAAAACTG TCGATGCTCA TCGCCTAGCA ATGAAAAAAT	1566
TTTTTCTTTT TTTTTCATT ATTTCTCTTT GTTGCGTACT TTTTTTCATT GCGTTTCGCG	1626
GCAAAAGCGA TTCGAGTTGA CTGGAAGTGT GTTATACTAT AAAAAGTGTA TATGCCTATT	1686
TTTGGTTCTG ATCTTTACTT TACTGTTAAG TACTGGCTGA GGCAGTAGAC TCTGCCTCTG	1746
TTACGGCAGC GGTATTCGCC TCGGCATCAG CAGCCGCCCA CGGTAGAGTA GGTTCTGTTG	1806
TTTTGACGTT TGCCAAGGTA CTGTCCAAAT GCTCCTTCAG CAAGGCCTCA TTACTTTCCT	1866
TCTCCGGACC CACCGATTGC GTGATCTCCT GTACACGGTT CAAGAACTTG TTCAAATTGT	1926
AGCCCGCAGC AGCATCAGAG ACTTCTTGTG TGTAAGGGAC ACCCCTCAAC TCCTTGACTC	1986
TTCTTTTGTG CACTTTGCCC TTTAAATGCG TTTTTAACGC TATAGCAGTC TCCATGTATT	2046
TGGCACAGTG TATGCAATAG TGCTGACCAA GGCCCGGTTT GGTTTCATCC AATGGCTGGT	2106
TCAGAAGCTT CTGTACTGAT TCCTTGGTGG ACAAATCGTT ATAGATCAGG TCCAAGTCTC	2166
GTGITCTTCT TTTAGTCTTG TATCTCTTCA CCGAATATCT ACCCATGATG CGCTATTGTT	2226
TTATCTTCAC TTGTCTGTGT GTTTAACTGC CTTTCAATTC ACCTCATCTC ATCTCCCGCT	2286
ACTITICCATA TATAAAAGCA AAATTAATIT GCTITITICCC CIGTCAGTAT AAAAAAATTT	2346
TCCGCAGGAT ATAGAAAAAA AAGAAATGAA ATTATAGTAG CGGTTATTTC CGTGGGGTGC	2406
TITTITACAC CIGTACATCT TITCCCTCCG TACATTITIT TTATTITIT TITGGGITIT	2466
TTTTTTCGA TATTTTTCCC TCCGAAACTA GTTAGCACAA TAATGCTGAC TAAGGAAACT	2526
TTTCATCTCA GAATTGATGG TCAGTTTGGT TTCTCTAGAG AATAGTTTAT AAAAAGATGT	2586
TGATGTGGAG CAACCATTTA TACATCCTTT CCGCAAGTGC TTTTGGAGTG GGACTTTCAA	2646
ACTITIANAGI ACAGIATATC AAATAACIAA TICAAGAIGG CIAGAAGACC AGCIAGAIGI	2706
FACAGATACC AAAAGAACAA GCCTTACCCA AAGTCTAGAT ACAACAGAGC TGTTCCAGAC	2766
CCCAAGATCA GAATCTACGA TTTGGGTAAG AAGAAGGCTA CCGTCGAT	2814

(2) INFORMATION FOR SEQ ID NO:43:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 14 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

### WHAT IS CLAIMED IS:

- A host cell transformed or transfected with DNA comprising:
  - a repressor gene encoding a repressor protein, said repressor gene under transcriptional control of a promoter;
  - a selectable marker gene encoding a selectable marker protein; said selectable marker gene under transcriptional control of an operator; said operator regulated by interaction with said repressor protein;
  - a first recombinant fusion protein gene encoding a first binding protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activating protein or a transactivating domain of a transcriptional activating protein; and
  - a second recombinant fusion protein gene encoding a second binding protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activating protein or a transactivating domain of a transcriptional activating protein, whichever domain is not encoded by the first fusion protein gene, said second binding protein or binding fragment thereof capable of interacting with said first binding protein or binding fragment thereof such that interaction of said second binding protein or binding fragment thereof and said first binding protein or binding fragment thereof brings into proximity a DNA binding domain and a transactivating domain forming a functional transcriptional activating protein; said functional transcriptional activating protein; said functional transcriptional activating protein said promoter to increase expression of said repressor gene.

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- The host cell of claims 1 wherein said DNA binding domain and said transactivating domain are derived from a common transcriptional activating protein.
- 3. The host cell of claim 1 wherein one or more of the repressor gene, the selectable marker gene, the first recombinant fusion protein gene, and the second recombinant fusion protein gene are encoded on distinct DNA expression constructs.
- 4. The host cell of claim 1 wherein said selectable marker protein is an enzyme in a pathway for synthesis of a nutritional requirement for said host cell such that expression of said selectable marker protein is required for growth of said host cell on media lacking said nutritional requirement.
- 5. The host cell of claim 1 wherein said host cell is a yeast cell or a mammalian.
- The host cell of claim 2 wherein said selectable marker gene encodes HIS3;
- 7. The host cell of claim 2 wherein said repressor protein gene encodes a tetracycline resistance protein;
- The host cell of claim 2 wherein said operator is a tet operator.
- 9. The host cell of claim 2 wherein said promoter is selected from the group consisting of the LexA promoter, the alcohol dehydrogenase promoter, the Gal4 promoter.

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- The host cell of claim 2 wherein said DNA binding domain derived from a protein selected from the group consisting of LexA and Gal4.
- The host cells of claim 2 wherein said transactivating domain is derived from a protein selected from the group consisting of VP16 and Gal4.
- The host cell of claim 2 wherein the first binding protein is CREB and the second binding protein is CBP.
- The host cell of claim 2 wherein the first binding protein is Tax and the second binding protein is SRF.
- 14. The host cell of claim 2 wherein the first binding protein is casein kinase I and the second binding protein is CREB.
- 15. The host cell of claim 2 wherein the first binding protein is AKAP 79 and the second binding protein is selected from the group consisting of RI, RII and calcineurin.
- 16. A method to identify an inhibitor of binding between a first binding protein or binding fragment thereof and a second binding protein or binding fragment thereof comprising the steps of:
  - a) growing host cells of any one of claims I through 15 in the absence of a test compound and under conditions which permit expression of said first binding protein or binding fragment thereof and said second binding protein or binding fragment thereof such that said first binding protein or fragment thereof and second binding protein or binding fragment thereof interact bringing

into proximity said DNA binding domain and said transactivating domain forming said functional transcriptional activating protein; said transcriptional activating protein acting on said promoter to increase expression of said repressor protein; said repressor protein interacting with said operator such that said selectable marker protein is not expressed;

- confirming lack of expression of said selectable marker protein in said host cell:
- growing said host cells in the presence of a test compound; and
- d) comparing expression of said selectable marker protein in the presence and absence of said test compound wherein increased expression of said selectable marker protein is indicative that the test compound is an inhibitor of binding between said first binding protein or binding fragment thereof and said second binding protein or binding fragment thereof.

# 17. The method of claim 16 wherein

the host cell is a yeast cell:

the selectable marker gene encodes HIS3:

transcription of the selectable marker gene is regulated by the *tet* operator;

the repressor protein gene encodes the tetracycline resistance protein:

transcription of the tetracycline resistance protein is regulated by the LexA promoter;

the DNA binding domain is derived from LexA; and the transactivating domain is derived from VP16. WO 98/13502 PCT/US97/17276

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18. The method of claim 16 wherein

the host cell is a yeast cell;

the selectable marker gene encodes HIS3;

transcription of the selectable marker gene is regulated by the *tet* operator;

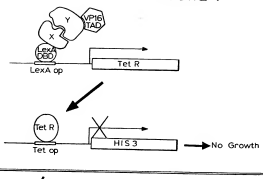
the repressor protein gene encodes the tetracycline resistance protein;

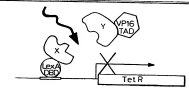
transcription of the tetracycline resistance protein is regulated by the alcohol dehydrogenase promoter;

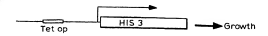
the DNA binding domain is derived from LexA; and the transactivating domain is derived from VP16.

19. A kit to identify an inhibitor of binding between a first binding protein or binding fragment thereof and a second binding protein or binding fragment thereof, said inhibitor identified by the method of claim 16.

FIGURE 1







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# INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (BCT)

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15/81, 1/19, C12Q 1/68	A3.	(43) International Publication Date: 2 April 1998 (02.04.98)
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(20) Priority Data: 08/721,730 27 September 1996 (27.09.9 (71) Applicant: ICOS CORPORATION [US/US): 22/ Avenue, S.E., Bothell, WA 98021 (US). (72) Inventors: GODDMAN, Richard, H.; 18560 Westvic Labe Oswego, OR 97034-7382 (US). HOEKSTR F, 10321 216th, S.E., Sonbomish, WA 98230 (US) (74) Agent: WILLIAMS, Joseph, A., Jr.; Marshall, Gentein, Murray & Bonn., 6300 Sears Tower Wacker Drive, Chicago, IL 60606-6402 (US).	D21 20 cw Driv IA, Me S).	(88) Date of publication of the international scarch report: 16 July 1998 (16.07.98) 1, 16.
(54) Title: .METHOD TO IDENTIFY COMPOUNDS FO	R DISI	. RUPTING PROTEIN/PROTEIN INTERACTIONS
(57) Abstract The present invention relates generally to materials and methods for identification of inhibitors of interactions between known binding partner proteins.		tera op Tet R  Tet op No Growth
		Tet R  Growth

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# INTERNATIONAL SEARCH REPORT

Intern sul Application No PCT/US 97/17276

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Special categories or  A' document defining considered to be E' earlier document filing date ' document which in obtation or other a 'document referring other means  C' document referring document referring there means  do	international filling date th the application but thereory underlying the a claimed invention a claimed invention a claimed invention decourant is taken alone alclaimed invention invention step when the more other auch does- floors to a person stillled in family servin report		
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# INTERNATIONAL SEARCH REPORT

national application No

PCT/US 97/17276

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	rnational Search Report has not been established in respect of certain claims under Article 17(2)(e) for the following reasons:
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2. 🗌	Claims Nos.: because they retale to parts of the international Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically:
3. 🗌	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(e).
Box ii	Observations where unity of invention is lacking (Continuation of Item 2 of first aheet)
This Inte	mational Searching Authority found multiple inventions in this international application, as follows:
	See annex
۵.	As all required additional sourch floss were timely paid by the applicant, this International Search Report covers all searchable claims.
2. 🗌	As all searchable claims could be searched without offort justifying an additional fee, this Authority did not invite payment of any additional fee.
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4. X	No required additional search fees were limity paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nes.:
	1-12, 16-19
-	*
Remark	on Protest  The additional search toos were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

### 1. Claims: 1-12, 16-19

A host cell transformed or transfected with DNA comprising: a repressor gene encoding a repressor protein, said repressor gene under transcriptional control of a promoter; a selectable maker gene encoding a selectable protein; said marker gene under transcriptional control of an operator; said operator regulated by interaction with said repressor protein; a first recombinant fusion protein gene encoding a first binding potein or binding fragment thereof in frame with either a transactivator domain of a transcriptional activator protein; and a second recombinant fusion protein or binding fragment thereof in frame with either a DNA binding domain of a transcriptional activating protein, whichever domain is not encoded by the first fusion protein gene, said second binding protein or binding fragment thereof capable of interacting with said first binding protein or binding fragment thereof such that interaction of said second binding protein or binding fragment thereof brings into proximity a DNA binding domain and a transactivating domain forming a functional transcriptional activating protein; said functional transcriptional activating protein acting on said promoter to increase expression of said repressor gene; said DNA binding domain and said transactivating domain are derived from a common transcriptional activating protein; one or more of the repressor gene, the selectable marker gene, and the first and second recombinant fusion protein genes, are encoded on distinct DNA expression constructs; said host cell wherein said selectable marker protein is an enzyme; said host cell is a yeast cell or a mammalian; said selectable marker gene encodes HIS3, said repressor protein gene encodes a tetracyline resistant protein; said operator is a tet operator; said promoter is selcted from the group consisting of the LexA-, the alcohol dehydrogenase-, the GA14-promoter; said DNA binding domain derived from a protein from the group consisting of LexA and Gal4; said transactivating domain is derived from a protein selected from the group consisting of VP16 and Gal4; said first binding protein is CREB and the second binding protein is CBP; a method and kit to identify an inhibitor of binding between a first and a second binding protein using said host cell.

### 2. Claim: 13

The host cell of subject one but wherein the first binding protein is Tax and the second binding protein is SRF.

### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

### 3. Claim: 14

The host cell of subject one but wherein the first binding protein is casein kinase I and the second binding protein is CREB.

### 4. Claim: 15

The host cell of subject one but wherein the first binding protein is AKAP 79 and the second binding protein is selected from the group consisting RI, RII and calcineurin.

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